IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

DECLARATION OF JOHN C. ROCKETT, Ph.D.

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- 1. Since 1995 I have been engaged full-time in molecular toxicology research, with an emphasis on the application of expression profiling techniques, including but not limited to nucleic acid microarray expression profiling techniques, to studies of the mechanisms of toxicant action and to the design of assays to monitor toxicant exposure.
- 2. My curriculum vitae, including my list of publications, is attached hereto as Exhibit A.
- 3. For the past 5 years, my work has focused primarily on analyzing the effects of potentially hazardous environmental agents, such as heat, water disinfectant byproducts, and conazole fungicides on the male reproductive tract. Although we are interested in the basic mechanisms of action of such toxicants, we also have two practical goals in mind: first, to identify individual agents and families of agents that adversely affect male reproductive development and function, and second, to develop methods for monitoring human exposure to such agents, particularly methods capable of identifying toxicant exposure at an early stage.
- 4. I have relied on expression profiling as a principal approach to these goals. Expression profiling, by

reporting the expression levels of thousands of genes simultaneously, gives us an opportunity to identify and group toxicants based on similarities in the patterns of gene expression they induce in cells and tissues; the gene expression profiles induced by treatment with known testicular toxins serve as standards; molecular signatures or molecular fingerprints as it were, against which the patterns of gene expression induced by agents of unknown toxicity may be compared and judged. In addition, gene expression profiling may give us the opportunity to detect toxicity before more gross phenotypic changes become manifest.

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served on the Microarray Technical
Subcommittee of the United States Environmental
Protection Agency (EPA) Genomics Task Force, and

served on the Scientific Committee for the conference series on "Critical Assessment of Techniques for Microarray Data Analysis," held annually at Duke University, Durham, NC;

and I currently

serve on the Technical Committee on the Application of Genomics to Mechanism-Based Risk Assessment of the International Life Sciences Institute's Health and Environmental Sciences Institute,

serve on the Genomics and Proteomics Committee of the National Health and Environmental Effects Research Laboratory of the EPA's Office of Research and Development,

belong to the [North Carolina Research]
Triangle Array Users Group,

the Molecular Biology file systems, it Speciality Section of the Society of Toxicology, and

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belong to the Triangle Consortium for Reproductive Biology 1996 Laborate the Continue of Manager that more research to their he arrest had completely

In addition, I am the principal investigator on a cooperative research and development agreement (CRADA) entitled "Development of a Genetic Test for Male Factor Infertility." Prior to this, I was a co-principal investigator on a ... materials cooperative research and development agreement (MCRADA) to print oligonucleotide-based microarrays; and from 1999 - 2002, I was coinvestigator on a CRADA to develop gene microarrays for toxicology applications. Selections

- I presume the reader's familiarity with the basic construction and operation of microarrays. For purposes of the discussion to follow, I use the phrase "nucleic acid microarray" and, equivalently, the term "microarray" to refer generically to the various types of nucleic acid microarray that include immobilized nucleic acid probes of sufficient length to permit specific binding, with minimal crosshybridization, to the probe's cognate transcript, whether the transcript is in the form of RNA or DNA. Although this definition excludes microarrays having shorter probes, such as the 20-mer probes of arrays manufactured by Affymetrix, Inc., many of the comments that follow nonetheless apply to such microarrays as well.
- Although my own work with microarrays dates back only to 1998, and high density spotted nucleic acid

microarrays themselves date back perhaps only to 1995, microarrays are by no means the only, nor the first, expression profiling tool. As I describe in detail in my Xenobiotica review, there are a number of other differential expression analysis technologies that precede the development of microarrays, some by decades, and that have been applied to drug metabolism and toxicology research, including:

(1) differential screening; (2) subtractive hybridization, including variants such as chemical cross-linking subtraction, suppression-PCR subtractive hybridization and representational difference analysis; (3) differential display; (4) restriction endonuclease facilitated analyses, including serial analysis, of gene expression (SAGE) and gene expression fingerprinting; and (5) EST analysis.

8. In my own earlier research, I used both reverse-transcriptase polymerase chain reaction (RT-PCR) and suppression-PCR subtractive hybridization (SSH) to study patterns of differential gene expression caused by hepatic challenge with nongenotoxic and genotoxic hepatotoxins.

Schena et al., "Quantitative monitoring of gene expression patterns with a complementary DNA microarray," Science 270:467-470 (1995), attached hereto as Exhibit B.

Rockett et al., "Differential gene expression in drug metabolism and toxicology: practicalities, problems and potential," Xenobiotica 29:655-691 (1999) (hereinafter, "Xenobiotica review"), attached hereto as Exhibit C.

See, e.g., Rockett et al., "Molecular profiling of non-genotoxic carcinogenesis using differential display reverse transcription polymerase chain reaction (ddRT-PCR)," European J. Drug Metabolism & Pharmacokinetics 22(4):329-33 (1997), and Rockett et al., "Use of a suppression-PCR subtractive hybridization method to identify gene species which demonstrate altered expression in male rat and guinea pig livers following 3-day exposure to [4-chloro-6-(2,3-xylidino)-2-pyrimidinylthio) acetic acid," Toxicology 144(1-3):13-29 (2000), attached hereto respectively as Exhibits D and E.

- These older transcript expression profiling techniques provide analogous expression data, but with far lower throughput.
- the introduction of high density spotted microarrays in 1995, that;

The other was considered as the the section of

- (i) each probe on the microarray, with careful design and sufficient length, and with sufficiently stringent hybridization and wash conditions, binds specifically and with minimal cross-hybridization, to the probe's cognate transcript;
 - (ii) each additional probe makes an additional transcript newly detectable by the microarray, increasing the detection range, and thus versatility, of this analytical device for gene expression profiling;
 - (iii) it is not necessary that the biological function be known in order for the gene,

The compelling logic of this proposition has likely motivated the remarkably rapid progress from the earliest high density spotted arrays in 1995 (Schena et al., "Quantitative monitoring of gene expression patterns with a complementary DNA microarray, * Science 270:467-470 (1995), attached hereto as Exhibit B), to the first whole genome arrays in 1997 (Lashkari et al., "Yeast microarrays for genome wide parallel genetic and gene expression analysis, Proc. Natl. Acad. Sci. USA 94(24):13057-62 (1997) and DeRisi et al., *Exploring the metabolic and genetic control of gene expression on a genomic scale, Science 278(5338):680-6 (1997), attached hereto as Exhibits F and G, respectively), to the concurrent announcement by two companies earlier this month of their respective commercial introductions of single chip human whole genome arrays (Pollack, *Human Genome Placed on Chip; Biotech Rivals Put it Up for Sale, " The New York Times, Thursday, October 2, 2003 (Business Day), attached hereto as Exhibit H; *Agilent Technologies ships whole human genome on single microarray to gene expression customers for evaluation, Press Release, Agilent Technologies, October 2, 2003, attached hereto as Exhibit I; "Affymetrix Announces Commercial Launch of Single Array for Human Genome Expression Analysis; More Than 1 Million Probes Analyze Expression Levels of Nearly 50,000 RNA Transcripts and Variants on a Single Array the Size of a Thumbnail, Press Release, Affymetrix, October 2, 2003, attached hereto as Exhibit J).

or a fragment of the gene, to prove useful as a probe on a microarray to be used for expression analysis;

- (iv) failure of a probe to detect changes in expression of its cognate gene does not diminish the usefulness of the probe on the microarray; and
- (iv) failure of a probe to detect a particular transcript in any single experiment does not deprive the probe of usefulness to the community of users who would use this research tool.

These principles also apply to transcript expression profiling techniques that antedate the development of high density spotted microarrays, and accordingly were well-understood prior to 1995.

- 11. Moreover, expression profiling is not limited to the measurement of mRNA transcript levels. It is widely understood among molecular and cellular biologists that protein expression levels provide complementary profiles for any given cell and cellular state. Although I cannot claim credit for having coined the phrase, I have written that the difference between transcript expression profiling and protein expression profiling is that "transcriptomics indicates what should happen, and proteomics shows what is happening." 5
- 12. For decades, such protein expression profiles have been generated using two dimensional polyacrylamide gel

Rockett, "Macroresults through Microarrays," Drug Discovery Today 7:804 - 805 (2002) (emphasis added), attached hereto as Exhibit K.

electrophoresis (2D-PAGE), and used, among other things, to study drug effects.

- 13. Although the protein expression profiles produced by 2D-PAGE analysis are analogous to the transcript expression profiles provided by nucleic acid microarrays, an even closer analogy is perhaps offered by antibody antibody microarrays; as I note in my Drug Discovery Today commentary, such antibody microarrays date back to the work of Roger Ekins in the mid- to late-1980s.7
- protein expression profiling analyses, particularly to analyses performed using antibody microarrays. Thus, as with nucleic acid microarrays, the greater the number of proteins detectable, the greater the power of the technique; the absence or failure of a protein to change in expression levels does not diminish the usefulness of the method; and prior knowledge of the biological function of the protein is not required. As applied to protein expression profiling, these principles have been well understood since at least as early as the 1980s.
- 15. Both gene and protein expression profiling are particularly useful to the toxicologist, especially in the pharmaceutical industry. Accordingly, I made the following

See, e.g., Anderson et al., "A two-dimensional gel database of rat liver proteins useful in gene regulation and drug effects studies," Electrophoresis 12:907 - 930 (1991), attached hereto as Exhibit L.

See Ekins et al., J. Bioluminescence Chemiluminescence 5:59-78 (1989); Ekins et al., Clin. Chem. 37: 1955-1965 (1991); and Ekins, U.S. Patent Nos. 5,432,099, 5,807,755, and 5,837,551, attached hereto respectively as Exhibits M to Q.

statements in my Xenobiotica review, written in the summer of 1998: Laboration of the summer of the

[I]n the field of chemical-induced toxicity, it is now becoming increasingly obvious that most adverse reactions to drugs and chemicals are the result of multiple gene regulation, some of which are causal and some of which are casually-related to the toxicological phenomenon per seing the servation has led to an upsurge in interest in gene-profiling technologies which differentiate between the control and toxin-treated gene pools in target tissues and is, therefore, of value in rationalizing the molecular mechanisms of exemploiotic-induced toxicity.

Knowledge of toxin-dependent gene regulation in target tissues is not solely an academic pursuit as much interest has been generated in the pharmaceutical industry to harness this technology in the early identification of toxic drug candidates, thereby shortening the developmental process and contributing substantially to the safety assessment of new drugs.

For example, if the gene profile in response to say a testicular toxin that has been well-characterized in vivo could be determined in the testis, then this profile would be representative of all new drug candidates which act via this specific molecular mechanism of toxicity, thereby providing a useful and coherent approach to the early detection of such toxicants.

Whereas it would be informative to know the identity and functionality of all genes up/down regulated by such toxicants, this would appear a longer term goal, as the majority of human genes have not yet been sequenced, far less their functionality determined. However, the current use of gene profiling yields a pattern of gene changes for a xenobiotic of unknown toxicity which may be matched to that of well-characterized toxins, thus alerting the toxicologist to possible in vivo similarities between the unknown and the standard.

Despite the development of multiple technological advances which have recently brought the field of gene expression profiling to the forefront of molecular analysis, recognition of the importance of differential gene expression and characterization of differentially expressed genes has existed for many years.

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16. As noted in the preceding excerpt from my Xenobiotica review, expression profiling in toxicology studies yield patterns of changes that are characteristic of an agent of unknown toxicity, which patterns may usefully be matched to those of well-characterized toxins.

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17. In the context of such patterns of gene expression, each additional gene-specific probe provides an additional signal that could not otherwise have been detected, giving a more comprehensive, robust, higher resolution -- and thus more useful -- pattern than otherwise would have been possible.

18. It is my opinion, therefore, based on the state of the art in toxicology at least since the mid-1990s -- and as regards protein profiling, even earlier -- that disclosure of the sequence of a new gene or protein, with or without knowledge of its biological function, would have been

⁸ In a sense, each gene-specific probe used in such an analysis is analogous to a different one of the many parts of an engine, with each individual part, or subcombinations of such parts, deriving at least part of their usefulness from the utility of the completed combination, the functioning engine.

sufficient information for a toxicologist to use the gene and/or protein in expression profiling studies in toxicology.

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- represent my individual views and are not intended to represent the opinion of my employer, the United States Environmental Protection Agency, or of any other branch of the federal government. Other than my current engagement to provide this declaration, I have neither had, nor currently have, financial ties to, or financial interest in, Incyte Corporation. I am not myself an inventor on any patent application claiming a gene or gene fragment.
- 20. I declare further that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true, and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and may jeopardize the validity of any patent application in which this declaration is filed or any patent that issues thereon.

John Coughlin Rockett III, Ph.D.

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Supervisors: Dr Sally P. Darney (Scientific publications under Sally D. Perreault) and Dr David and the state of t J. Dix.

5/95-7/98: Rhone-Poulenc Post-Doctoral Research Fellow, Molecular Toxicology Group, School of Biological Sciences, University of Surrey, Guildford, Surrey, England. "Supervisor: Prof. G. Gordon Gibson. 16. 4071 (1994) 1994

the control of the co Ph.D., 1995 - University of Warwick, Coventry, W. Midlands, England

Title: Transforming Growth Factor-β and Immune Recognition Molecules in Oesophageal

Supervisors: Dr Alan G. Morris (University of Warwick) and Dr S. Jane Darnton (Birmingham Heartlands Hospital)

B.Sc. (Hons.), 1991 - University of Warwick, Coventry, W. Midlands, England. Degree: Microbiology and Microbial Technology (with intercalated year in industry), Class 2i. Tutor: Professor Howard Dalton.

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Society of Toxicology (Inc. Molecular Biology Speciality Section) (2001-present) Science Advisory Board (2001-present)

North Carolina Chapter of the Society of Toxicology (1999-present)

Triangle Consortium for Reproductive Biology (1999-present)

Biology of Toxicology (1999-present)

Institute of Biology (U.K.) (1989 - present)

British Toxicology Society (1996 - 2000)

Biochemical Society (U.K.) (1992-1995)

British Society for Immunology (1992-1995) Susteen him and the Applied Tokic dov Entish Society for immunology (1992-1993)

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Membership of Scientific Committees:

International Life Sciences Institute's (ILSI) Health and Environmental Sciences Institute (HESI) Technical Committee on the Application of Genomics to Mechanism-Based Risk Assessment:

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• Steering Committee (5/02-present).

Hepatotoxicity Working Group Vice-Chair (5/02-present).

Hepatotoxicity Work Group Member (5/01-present).

Charter member, Fertility and Early Pregnancy Work Group of the National Children's Study (07/01-Present).

National Health and Environmental Effects Research Laboratory Distinguished Lecture Series Committee (July 03-present).

U.S. Environmental Protection Agency Genomics Task Force Microarray Technical Subcommittee (August 03-present).

National Health and Environmental Effects Research Laboratory Genomics and Proteomics Committee (NGPC) (September 03-present).

Professional Meetings:

Invited participant ("Observer") in Expert Panel Workshop: "The Role of Environmental Factors on the Onset and Progression of Puberty in Children". Organised by Serono Symposia International. November 6th-8th, 2003, Chicago, IL, USA.

Joint organiser and co-chair of: "Genomic analysis of surrogate tissues for measuring toxic exposures and drug action", the "Innovations in Applied Toxicology" Symposium for the Society of Toxicology 42nd Annual Meeting, March 9th-13th, 2003, Salt Lake City, UT, USA.

- (8) John C. Rockett, David J. Esdaile and G Gordon Gibson (1999). Differential gene expression in drug metabolism: practicalities, problems and potential. Xenobiotica, 29(7):655-691.

 (7) MC Murphy, CN Brookes, JC Rockett, C Chapman, JA Lovegrove, BJ Gould, JW Wright and CM Williams (1999). The quantitation of lipoprotein lipase mRNA in biopsies of human adipose tissue, using the polymerase chain reaction, and the effect of increased consumption of n-3 polyunsaturated fatty acids. European Journal of Clinical Nutrition, 53:441-447.
- (6) JC Rockett, DJ Esdaile and GG Gibson (1997). Molecular profiling of non-genotoxic carcinogenesis using differential display reverse transcription polymerase chain reaction (ddRT-PCR). European Journal of Drug Metabolism & Pharmacokinetics 22(4):329-33.
- (5) Rockett, J., Larkin, K., Darnton, S., Morris, A. and Matthews, H. (1997). Five newly established oesophageal carcinoma cell lines: phenotypic and immunological characterisation. *British Journal of Cancer* 75(2):258-263.
- (4) J C Rockett, S J Darnton, J Crocker, H R Matthews and A G Morris (1996). Lymphocyte infiltration in oesophageal carcinoma: lack of correlation with MHC antigens, ICAM-1, and tumour stage and grade. Journal of Clinical Pathology 49:264-267.
- (3) J C Rockett, S J Darnton, J Crocker, H R Matthews and A G Morris (1995). Expression of HL-ABC and HLA-DR histocompatability antigens and intercellular adhesion molecule-1 in oesophageal carcinoma. *Journal of Clinical Pathology* 48:539-44.
- (2) Salam M, Rockett J and Morris A (1995). The prevalence of different human papillomavirus types and p53 mutations in laryngeal carcinomas: is there a reciprocal relationship? European Journal of Surgical Oncology 21:290-296.
- (1) Salam M, Rockett J and Morris A (1995). General primer-mediated polymerase chain reaction for simultaneous detection and typing of HPV in laryngeal carcinomas. Clinical Otolaryngology 20:84-88.

(2) Articles Submitted To A Scientific Journal

- (4) John C. Rockett, Judith E. Schmid, Christopher J. Luft, J. Brian Garges, M. Stacey Ricci, Pasquale Patrizio, Norman B. Hecht and David J. Dix. Gene Expression Patterns Associated with Infertility in Rodent and Human Models. *An invited submission*
- (3) Roger Ulrich, John C. Rockett, G. Gordon Gibson and Syril Pettit. Evaluating the Effects of Methapyrilene and Clofibrate on Hepatic Gene Expression: A Collaboration Between Laboratories and a Comparison of Platform and Analytical Approaches.
- (2) Valerie A Baker, Helen M Harries, Jeffrey F Waring, Roger Jolly, Angus de Souza, Judith E Schmid, Hong Ni, Roger Brown, Roger G Ulrich and John C. Rockett. Clofibrate-Induced Gene Expression Changes in Rat Liver: A Cross-Laboratory Analysis Using Membrane cDNA Arrays.

(1) David Miller, Corrado Spadafora, David Dix, Adrian Platts, John C. Rockett, Stephen A Krawetz Nuclease digestion of sperm chromatin suggests a random distribution of gene sequences.

(3) Articles In Preparation For Submission To A Scientific Journal

- (3) Spearow J, DB Tully, JC Rockett and DJ Dix. Differential testicular gene expression in mouse strains sensitive and resistant to endocrine disruption by estrogen.
- (2) Sally D. Perrault, John C. Rockett, Laura Fenster, James Kesner, Wendy Robbins and Steven Schrader. Biomarkers for Assessing Reproductive Development and Health: Part 2 Adult Reproductive Health.
- (1) J. Christopher Luft, Douglas B. Tully, John C. Rockett, Judith E. Schmid and David J. Dix. Reproductive and genomic effects in testes from mice exposed to the water disinfectant byproduct bromochloroacetic acid

(4) Book Chapters

- (4) John C. Rockett. Gene Microarrays Applied to Reproductive Toxicology. In Cunningham (Ed): Genetic and Proteomic Applications in Toxicity Testing, The Human Press, Totowa. In Preparation. *An invited submission*
- (3) John C. Rockett and David J Dix. Gene Expression Networks. In Cooper (ed-in-chief): Encyclopaedia of the Human Genome, Nature Publishing Group. London, New York. ISBN 0-333-80386-8 (2003). *An invited submission*
- (2) John C. Rockett. The Future of Toxicogenomics. In Michael Burczynski (ed): "An Introduction to Toxicogenomics". CRC Press. Boca Raton, London, New York, Washington D.C., pp299-317 (2003). *An invited submission*
- (1) J. Rockett, S. Darnton, J. Crocker, H. Matthews and A. Morris: Major Histocompatibility Complex (MHC) class I and II and Intercellular Adhesion Molecule (ICAM)-1 expression in oesophageal carcinoma. Peracchia A, Rosati R, Bonavina L, Bona S, Chella B (eds): Recent Advances in Diseases of the Esophagus. Bologna: Monduzzi Editore, pp45-49 (1996).
- (5) Other Scientific Publications (Letters to Editors; Meeting Reports; Commentaries etc.)
 - (11) John C. Rockett (2003). Probing the nature of microarray-based oligonucleotides. Drug Discovery Today 8(9):389. (A Letter To The Editor) *An invited submission*
 - (10) John C. Rockett (2003). To confirm or not to confirm (microarray data) that, is the question. Drug Discovery Today 8(8):343. (A Letter To The Editor)

- (9B) Nazzareno Ballatori, James L. Boyer, and John C. Rockett. (2003), Exploiting Genome Data to Understand the Function, Regulation and Evolutionary Origins of Toxicologically Relevant Genes. Environ Health Perspect. 111(6):871-5. (A Meeting Report)
- (9A) Nazzareno Ballatori, James L. Boyer, and John C. Rockett. (2003). Exploiting Genome Data to Understand the Function, Regulation and Evolutionary Origins of Toxicologically Relevant Genes. EHP Toxicogenomics. 111(1T):61-5. (A Meeting Report)
- (8) John C. Rockett (2002). Surrogate Tissue Analysis for Monitoring the Degree and Impact of Exposures in Agricultural Workers. AgBiotechNet, 4:1-7 November, ABN 100. (A Review Article). *An invited submission*
- (7) John C. Rockett (2002). Macroresults Through Microarrays. Drug Discovery Today, 7(15);804-805. (A Meeting Report)
- (6) John C. Rockett (2002). Chip, chip, array! Three chips for post-genomic research. Drug Discovery Today, 7(8);458-459. (A Meeting Report)
- (5) John C. Rockett (2002). Use of Genomic Data in Risk Assessment. GenomeBiology, 3(4): reports4011.1-4011.3 (http://genomebiology.com/2002/3/4/reports/4011/?isguard=1). (A Meeting Report)
- (4) John C. Rockett (2001). Genomic and Proteomic Techniques Applied to Reproductive Biology. GenomeBiology 2(9): 4020.1-4020.3 (http://genomebiology.com/2001/2/9/reports/4020/). (A Meeting Report)
- (3) John C. Rockett (2001). Chipping away at the mystery of drug responses. The Pharmacogenomics Journal, 1(3);161-163. (A commentary) *An invited submission*
- (2) Rockett, John C. and Dix, David J. (1999). U.S. EPA workshop: Application of DNA arrays to Toxicology. *Environmental Health Perspectives*, 107(8):681-685. (A Meeting Report)
- (1) John C. Rockett III (1995). Immune recognition molecules and transforming growth factor beta-1 in oesophageal cancer. Ph.D. thesis, University of Warwick, Coventry, England. (Ph.D. thesis)

(6) Published Book, Paper and Website reviews

(9) John C. Rockett (2002). A report on the manuscript: Systemic RNAi in C. elegans requires the putative transmembrane protein SID-1. Winston WM, Molodowitch C, Hunter CP. Science. 2002 295:2456-2459. GenomeBiology, 3(7):reports0034 http://genomebiology.com/2002/3/7/reports/0034/

- (8) John C. Rockett (2001). A report on the manuscript: Genetic rescue of an endangered mammal by cross-species nuclear transfer using post-mortem somatic cells. P Loi, et al., Nat Biotechnol. (2001, 19:962-964. GenomeBiology, 3(1):reports/0006/). (http://genomebiology.com/2001/3/1/reports/0006/).
- (7) John C. Rockett (2001). A report on the manuscript: Molecular Classification of Human Carcinomas by Use of Gene Expression Signatures. A Su et al., Cancer Res. 2001.61:7388-7393.

 GenomeBiology, 3(1):reports0005. (http://genomebiology.com/2001/3/1/reports/0005/).
- (6) John C. Rockett (2001). A report on the manuscript: Genetic evidence for two species of elephant in Africa. A Roca et al., Science. 2001 Aug 24;293(5534):1473-7. GenomeBiology, 2(12):reports0045. (http://www.genomebiology.com/2001/2/12/reports/0045/...
- (5) John C. Rockett (2001). A report on the manuscript: Extensive genetic polymorphism in the human CYP2B6 gene with impact on expression and function in human liver. T Lang et al., *Pharmacogenetics*, 2001, 11(5):399-415. *GenomeBiology*, 2(12):reports0044. (http://www.genomebiology.com/2001/2/12/reports/0044/).
 - (4) John C. Rockett (2001). A report on the manuscript: Novel Human Testis-Specific cDNA: molecular Cloning, Expression and Immunological Effects of the Recombinant Protein. R Santhanam and R K Naz, Molecular Reproduction and Development 60:1-12 (2001): GenomeBiology, 2(11):reports0040. (http://genomebiology.com/2001/2/11/reports/0040/).
 - (3) John C. Rockett (2001). A report on the website: BIND The Biomolecular Interaction Network Database (http://www.bind.ca/). GenomeBiology, 2(9): reports2011. http://www.genomebiology.com/2001/2/9/reports/2011/.
- (2) John C. Rockett (2001). A report on the manuscript: Exploring the DNA-binding specificities of zinc fingers with DNA microarrays. ML Bulyk et al., *Proc Natl Acad Sci USA* 2001, 98:7158-7163. GenomeBiology, 2(10): reports0032. (http://genomebiology.com/2001/2/10/reports/0032/).
 - (1) J Rockett (1996). A Book Review on: "Cell Adhesion and Cancer" (Eds., Hogg N. and Hart I.). Clinical Molecular Pathology 49(1):M64. *An invited submission*

(7) Published Abstracts of Poster and Oral Presentations

- (17) Amber K. Goetz, Wenjun Bao, Judith E. Schmid, Carmen Wood, Hongzu Ren, Deborah S. Best, Rachel N. Murrell, John C. Rockett, Michael G. Narotsky, Douglas C. Wolf, Douglas B. Tully, David J. Dix Gene Expression Profiling in Testis and Liver of Mice to Identify Modes of Action of Conazole Toxicities. Society of Toxicology 43rd Annual Meeting, March 21st-25th, 2004, Baltimore, MD, USA. *Toxicological Sciences*. (Submitted)
- (16) Jane Gallagher, Theresa Lehman, Ramakrishna Modali, Scott Rhoney, Marien Clas, Jeff Inmon, John C. Rockett, David Dix, Cindy Mamay, Suzanne Fenton, Suzanne McMaster, Stan

- Barone Jr, Pauline Mendola and Reeder Sams. Validation of Non-Invasive Biological Samples:

 'Pilot Projects Relevant to the National Children Study. Society of Toxicology 43rd Annual Meeting,

 March 21st 25th, 2004, Baltimore, MD, USA. *Toxicological Sciences*. (Submitted)
- (15) B.S. Pukazhenthi, J. C. Rockett, M. Ouyang, D.J. Dix, J.G. Howard, P. Georgopoulos, W.J. J. Welsh and D. E. Wildt. Gene Expression In The Testis Of Normospermic Versus Teratospermic Domestic Cats Using Human cDNA Microarray Analyses. Society for the Study of Reproduction 36th Annual Meeting, July 19th-22nd, 2003, Cincinnati, OH, USA. Biology of Reproduction 68 (Supp 1):191.
 - (14) David J. Dix and John C. Rockett (2003). Genomic and Proteomic Analysis of Surrogate Tissues for Assessing Toxic Exposures and Disease States. Innovation in Applied Toxicology symposium entitled "Genomic and Proteomic Analysis of Surrogate Tissues for Assessing Toxic Exposures and Disease States". Society of Toxicology 42nd Annual Meeting, March 9th-13th, 2003, Salt Lake City, UT, USA. Toxicological Sciences 72(S-1):276.
- (13) John C. Rockett, Chad R. Blystone, Amber K. Goetz, Rachel N. Murrell, Judith E. Schmid and David J. Dix. (2003). Gene Expression Profiling Of Accessible Surrogate Tissues To Monitor Molecular Changes In Inaccessible Target Tissues Following Toxicant Exposure. Innovations in Applied Toxicology Symposium entitled "Genomic and Proteomic Analysis of Surrogate Tissues for Assessing Toxic Exposures and Disease States". Society of Toxicology 42nd Annual Meeting, March 9th-13th, 2003, Salt Lake City, UT, USA. Toxicological Sciences 72(S-1):276.
- (12) Douglas B. Tully, J. Christopher Luft, John C. Rockett, Judy E. Schmid and David J. Dix (2002). Effects on gene expression in testes from adult male mice exposed to the water disinfectant byproduct bromochloroacetic acid. Society for the Study of Reproduction 35th Annual Meeting, July 28-31, 2002, Baltimore, Maryland, USA. Biology of Reproduction 66 (Supp 1):223.
- (11) David J. Dix, Kary E. Thompson, John C. Rockett, Judith E. Schmid, Robert J. Goodrich, David Miller, G. Charles Ostermeier and Stephen A. Krawetz (2002). Testis and spermatazola RNA profiles of normal fertile men. Society for the Study of Reproduction 35th Annual Meeting, July 28-31, 2002, Baltimore, Maryland, USA. Biology of Reproduction 66 (Supp 1):194.
- (10) Asa J. Oudes, **John C. Rockett**, David J. Dix and Kwan Hee Kim (2002). Identification of retinoic acid induced genes in mouse testis by cDNA microarray analysis. 27th Annual Meeting of the American Society of Andrology, 4/24-27/02. J. Andrology Supplement March/April.
- (9) John C. Rockett, Robert J. Kavlock, Christy Lambright, Louise G. Parks, Judith E. Schmid, Vickie S. Wilson and David J. Dix (2002). Use of DNA arrays to monitor gene expression in blood and uterus from Long-Evans rats following 17-β-estradiol exposure a new approach to biomonitoring endocrine disrupting chemicals using surrogate tissues. Toxicological Sciences 66(1): Abstract No.1388.
- (8) David J. Dix and John C. Rockett (2002). Genomic analysis of the testicular toxicity of haloacetic acids. Platform presentation at the symposium, "Defining the cellular and molecular

mechanisms of toxicant action in the testis". Toxicological Science 66 (1): Abstract No.848.

- (7) JC Rockett, JC Luft, JB Garges and DJ Dix (2001). The reproductive effects of the water disinfectant byproduct bromochloroacetate on juvenile and adult male mice. *Toxicological Sciences*, 60 (1):250.
- (6) Tarka DK, Klinefelter GR, Rockett JC, Suarez JD, Roberts NL and Rogers JM (2001). Effect of gestational expsore to ethane dimethane sulfonate (EDS), bromochloroacetic acid (BCA) and molinate on reproductive function in CD-1 male mice. *Toxicological Sciences*, 60 (1):250.
- (5) Garges JB, Rockett JC and Dix DJ (2001). Developmental and reproductive phenotype of mice lacking stress-inducible 70 kDa heat shock proteins (Hsp70s). *Toxicological Sciences*, 60 (1):383. (4) D Dix, J Rockett, J Luft, J Garges, M Ricci, P Patrizio and N Hecht (2000). Using DNA microarrays to characterise gene expression in testes of fertile and infertile humans and mice. *Biology of Reproduction*, 62 (s1);227.
- (3) J Luft, J B Garges, J Rockett and D Dix (2000). Male reproductive toxicity of bromochloroacetic acid in mice. *Biology of Reproduction*, 62 (s1);246.
- (2) Rockett, JC, Garges, JB and Dix, DJ (2000). A single heat-shock of juvenile male mice causes a long-term decrease in fertility and reduces embryo quality. *Toxicological Sciences* 54 (1):365.
- (1) JC Rockett, SJ Darnton, J Crocker, HR Matthews and AG Morris (1994). Major Histocompatability (MHC) class I and II and intercellular adhesion molecule (ICAM)-1 expression in oesophageal carcinoma (OC). Immunology 83 (s1):64.

(8) Invited Oral Presentations

- (10) John C. Rockett and Gary M Hellmann. To confirm or not to confirm (microarray data) that is the question. Seminar for EPA/NHEERL Genomics and Proteomics Committee's ArrayQA forum, August 25th, 2003, RTP, NC, USA.
- (9) John C. Rockett. "Biomonitoring Toxicant Exposure and Effect Using Toxicogenomics and Surrogate Tissue Analysis". Seminar for Division of Epidemiology, Statistics and Prevention Research, National Institute of Child Health and Development, May 29th, 2003, Rockville, MD, USA.
- (8) John C. Rockett. "Genomics and Proetomics: New Toxicity Testing". Platform presentation at US EPA Regional Risk Assessors Annual Conference, April 28th May 2nd, 2003, Stone Mountain, GA, USA.
- (7) John C. Rockett, Chad R. Blystone, Amber K. Goetz, Rachel N. Murrell, Judith E. Schmid and David J. Dix. "Gene Expression Profiling Of Accessible Surrogate Tissues To Monitor Molecular Changes in Inaccessible Target Tissues Following Toxicant Exposure." Platform presentation at

- SoT 42nd Annual Meeting symposium entitled "Genomic and Proteomic Analysis of Surrogate Tissues for Measuring Toxic Exposures and Drug Action", March 9th-13th, 2003, Salt Lake City, UT, USA.
- (6) John C. Rockett. "A Toxicogenomic Approach to Surrogate Tissue Analysis". Seminar for Department of Environmental and Molecular Toxicology, North Carolina State University, September 3rd, 2002, Raleigh, NC, USA.
- (5) John C. Rockett. "Differential gene expression in toxicology: practicalities, problems and potential". Platform presentation at 9th Annual Mount Desert Island Biological Laboratory Environmental Health Sciences Symposium: Exploiting Genome Data to Understand the Function, Regulation and Evolutionary Origins of Toxicologically Relevant Genes, July 10th-11th, 2002, Salisbury Cove, Maine, USA.
- (4) John C. Rockett, Leroy Folmar, Michael J. Hemmer and David J. Dix. "Arrays for biomonitoring environmental and reproductive toxicology". Platform Presentation at *Macroresults Through Microarrays 3 Advancing Drug Development*, April 29th-May 1st, 2002, Boston, MA, USA.
- (3) John C. Rockett, Sigmund Degitz, Suzanne E. Fenton, Leroy Folmar, Michael J. Hemmer, Joe E Tietge, and David J. Dix. "Use of DNA Arrays in Environmental Toxicology". Platform presentation at the 4th Annual Lab-on-a-Chip and Microarrays for Post-Genomic Applications meeting, January 14th-16th, 2002, Zurich, Switzerland.
- (2) John C. Rockett. "DNA Arrays". Seminar at EPA Molecular Biology Course, April 8th, 1999, USEPA, RTP, NC, USA.
- (1) John C. Rockett. "Contract Services for Array Applications". Seminar at the Triangle Array Users Group, May 1st, 1999, CIIT, RTP, NC, USA.

(9) Other Poster and Oral Presentations

- (23) John C. Rockett, Wenjun Bao, Chad R. Blystone, Amber K. Goetz, Rachel N. Murrell, Hongzu Ren, Judith E. Schmid, Jessica Stapelfeldt, Lillian F. Strader, Kary E. Thompson and David J. Dix. Genomic Analysis of Surrogate Tissues for Assessing Environmental Exposures and Future Disease States. ILSI-HESI meeting: Toxicogenomics in Risk Assessment Assessing the Utility, Challenges, and Next Steps. June 5th-6th, 2003, Fairfax, VA, USA.
- (22) John C. Rockett, Wenjun Bao, Chad R. Blystone, Amber K. Goetz, Rachel N. Murrell, Hongzu Ren, Judith E. Schmid, Jessica Stapelfeldt, Lillian F. Strader, Kary E. Thompson and David J. Dix. Genomic Analysis of Surrogate Tissues for Assessing Environmental Exposures and Future Disease States. EPA Science Forum, May 5th-7th, 2003, Washington, D.C., USA.

- (21) Germaine Buck, Courtney Johnson, Joseph Stanford, Anne Sweeney, Laura Schieve, John Rockett, Sherry Selevan and Steve Schrader. Prospective Pregnancy Study Designs for Assessing Reproductive and Developmental Toxicants. American Epidemiology Society Meeting, March 27th-28th, 2003, Atlanta, GA, USA.
- (20) John C. Rockett, Chad R. Blystone, Amber K. Goetz, Rachel N. Murrell, Hongzu Ren, Judith E. Schmid, Jessica Stapelfeldt, Lillian F. Strader, Kary E. Thompson, Doug B. Tully, Paul Zigas and David J. Dix. Genomic Analysis of Surrogate Tissues for Assessing Environmental Exposures and Future Disease States. *National Children's Study Assembly Meeting*, December 16th-18th, 2002, Baltimore, MD, USA.
- (19) John Rockett. The Use of Gene Expression Profiling to Detect Early Biomarkers of Adverse Effects Prior to Clinical manifestation. *National Children's Study: Meeting of EPA Project Leaders* Methods Development Projects. November 20th, 2002, USEPA, RTP, NC, USA. (Oral Presentation)
- (18) GC Ostermeier, RJ Goodrich, K Thompson, J Rockett, MP Diamond, K Collins, NICHD Reproductive Medicine Network, DJ. Dix, D Miller and SA Krawetz. Defining the spermatozoal RNA population in normal fertile men. American Society of Reproductive Medicine October 12-17, 2002, Seattle, WA, USA.
- (17) G. Charles Ostermeier, Robert J. Goodrich, Kary Thompson, John Rockett, Michael P. Diamond, Karen Collins, NICHD Reproductive Medicine Network, David J. Dix, David Miller and Stephen A. Krawetz. RNAs isolated from ejaculate spermatozoa provide a noninvasive means to investigate testicular gene expression. Gordon Conference on Mammalian Gametogenesis & Embryogenesis, June 30th-July 5th, Connecticut College, New London, CT, USA.
- (16) David Dix, John Rockett, Judith Schmid, Lillian Strader, Douglas Tully. Genomic analysis of testicular toxicity. *USEPA/NHEERL/RTD Peer Review*, October 22nd, 2001, RTP, NC, USA.
- (15) David Dix, John Rockett, Judith Schmid, Douglas Tully. Monitoring human reproductive health and development through gene expression profiling. *USEPA/NHEERL/RTD Peer Review*, October 22nd, 2001, RTP, NC, USA.
- (14) Patrizio P, N Hecht, J Rockett, J Schmid and D Dix (2001). DNA microarrays to study gene expression profiles in testis of fertile and infertile men. 57th Annual Meeting of the American Society for Reproductive Medicine, October 20th-25th, 2001, Orlando, FL, USA.
- (13) Jimmy L. Spearow, Dale Morris, Uland Wong, Rashid Altafi, Saeed Eteiwi, Mark Stanford, Trevor Stearns, Lorena Orozio, Angela Chen, John Rockett, Douglas Tully, David Dix and Marylynn Barkley. Genetic Variation In Susceptibility To The Disruption Of Testicular Development And Gene Expression By Pubertal Exposure To Estrogenic Agents. Third Annual University of California at Davis Conference for Environmental Health Scientists, Disruption of Developing Systems and Advances in Therapeutic Approaches August 27th, 2001, UC Davis, CA, USA.

- (12) Tarka DK, Klinefelter GR, Rockett JC, Suarez JD, Roberts NL and Rogers JM (2001). Effect of gestational expsore to ethane dimethane sulfonate (EDS), bromochloroacetic acid (BCA) and molinate on reproductive function in CD-1 male mice. North Carolina Society of Toxicology Winter Meeting, March 3rd, 2001. NIEHS, RTP, NC, USA.
- (11) David Dix, John Rockett, Leroy Folmar, Michael Hemmer, Sigmund Degitz, and Joseph Tietge (2001). Biomonitoring the Toxicogenomic Response to Endocrine Disrupting Chemicals in Humans, Laboratory Species and Wildlife. U.S. Japan International Workshop for Endocrine Disrupting Chemicals, February 28th-March 3rd, 2001, Tsukuba, Japan.
- (10) John C. Rockett, Faye L. Mapp, J. Brian Garges, J. Christopher Luft, Chisato Mori and David J Dix David Dix (2001). The effects of hyperthermia on spermatogenesis, apoptosis, gene expression and fertility in adult male mice. *Triangle Consortium for Reproductive Biology Annual Meeting*, January 27th, 2001, RTP, NC, USA.
- (9) Gangolli E, Dix DJ, Garges JB, Rockett, JC and Idzerda RL (2000). Testosterone Regulation of Sertoli Cell genes. 11th International Congress of Endocrinology, October 29th-November 2nd, 2000, Sydney, Australia.
- (8) J Rockett, J Luft, J Garges, M Ricci, P Patrizio, N Hecht and D Dix (2000). Using DNA microarrays to characterise gene expression in testes of fertile and infertile humans and mice. Functional Genomics & Microarray Data Mining, August 3rd-4thth 2000, Durham, NC, USA.
- (7) Rockett JC, S Ricci, P Patrizio, NB Hecht, JB Garges and DJ Dix (2000). Gene Expression in the Mammalian Testis. 5th NHEERL Symposium, June 6th-8th, 2000, RTP, NC, USA.
- (6) J Luft, J B Garges, J Rockett and D Dix (2000). Male reproductive toxicity of bromochloroacetic acid in mice. 2000 NIEHS/NTA Biomedical Science and Career Fair, April 28th 2000, RTP, NC, USA.
- (5) Rockett JC, S Ricci, P Patrizio, NB Hecht, JB Garges and DJ Dix (2000). Gene Expression in the Mammalian Testis. *Molecular Toxicology, Toxicogenomics and Associated Bioinformatics Applied to Drug Discovery* meeting, January 11th-15th, 2000, Santa Fe, NM, USA.
- (4) JC Rockett and DJ Dix (1999). Development of DNA arrays for the analysis of testis-expressed genes in humans and mice. The 8th Annual National Health and Environmental Effects Research Laboratory Open House. November 2nd-3rd, 1999, RTP, NC, USA.
- (3) **JC Rockett**, DJ Esdaile and GG Gibson (1997). Molecular profiling of non-genotoxic carcinogenesis using differential display reverse transcription polymerase chain reaction (ddRT-PCR). *The British Toxicology Society Annual Meeting*, April 19th-22nd, 1998, University of Surrey, Guildford, Surrey, England.
- (2) JC Rockett, DJ Esdaile and GG Gibson (1997). Molecular profiling of non-genotoxic

carcinogenesis using differential display reverse transcription polymerase chain reaction (ddRT-PCR). Poster presentation at Symposium on Drug Metabolism: Towards the next Millennium. August 26th-28th,1997, London King's College, London, England.

(1) J Rockett, S Darnton, J Crocker, H Matthews and A Morris: Major Histocompatibility Complex (MHC) class I and II and Intercellular Adhesion Molecule (ICAM)-1 expression in oesophageal carcinoma. Oral presentation at *The 6th World Congress of the International Society for Diseases of the Esophagus*, August 23rd-26th, 1995, Milan, Italy.

Artip sequence following Serene and coours within the domain of Artip that shows increading with HEE (14). To delete the complete STE23 acquence and create the state of the control of t TCTTGCTCATTTTGATATTGCTC- TGTAGATTG-TACTGAGAGTECAC-3"; and 5'-GCTACAAACAGC. GTCGACTTGAATGCCCCGACATCTTCGACTGT: GOGGTATTTCACACOG-31) were used to employ, the URAS sequence of pRSS16; and the reaction product was transformed into yearst for one-step gene replacement [R. Rolfistein, Methods Braymol. 194, 281 (1991). To create the aut A: LEU2 mutation contained on p114, a 5.0-kb Sal I tragment from pAXL1. is cloned into pUC19, and an internal 4.0-kb Hos i-Who I tragment was replaced with a LEU2 tragment. To construct the sta23A-LET2 affeit (a delation conresponding to 931 amino acids) carried on p153, a LEU2 tragment was used to reptace the 2.8-lib Pml I-Ec136 Il tragment of STE23, which occurs within a 6.24b Hind BI-Bd B genomic tragment cented on pSP72 (Promege). To create YEpMFA1, a 1.84b Barn HI tragment containing MFA1, from pIOC16 IC. Kuchler, R. E. Sterne, J. Thomer, EMBO J. 8, 3973 (1989)], was ligated into the Barn HI site of YEp351 [J. E. Hill, A. M. Myers, T. J. Koerner, A. Tzagoloff, Yeast 2, 163 (1986)

24. J. Chant and L. Herskowitz, Cell 65, 1203 (1991). 25. B. W. Matthews, Acc. Chem. Res. 21, 333 (1988).

26. K. Kuchier, H. G. Dohimen, J. Thomer, J. Cell Blot. 120, 1203 (1993); R. Koling and C. P. Hollenberg, EMBO J. 13, 3261 (1994); C. Berkower, D. Loeyza, S. Michaelis, Mcl. Biol. Cell 5, 1185 (1994).

- S. Michaels, Mol. Biol. Cell 5, 1180 (1994).

 27. A. Bender and J. R. Pringle, Proc. Natl. Acad. Sci.
 U.S. A 86, 9976 (1989); J. Chant, K. Corrado, J. R.
 Pringle, L. Herskowitz, Cell 65, 1213 (1991); S.
 Powers, E. Gonzales, T. Christensen, J. Cubert, D.
 Broek, Ibid., p. 1225; H. O. Park, J. Chant, I. Herskowitz, Nature 365, 269 (1993); J. Chant, Trands
 Genal. 10, 328 (1994). Genet. 10, 328 (1994);__ Genet. 10, 328 (1994); _____ and J. R. Pringle, J. Cell Biol. 129, 751 (1995); J. Chant, M. Mischke, E. Mitchell, L. Henskowitz, J. R. Pringle, bid., p. 767.
- 28. G. F. Sprague Jr., Methods. Enzymol. 194, 77 (1991).
- 29. Single-letter abbreviations for the amino acid resi-Class are as follows: A, Ale: C, Oys; D, Asp; E, Glu; F; Phe: G, Gly; H, Hie; L, Ie; K, Lys; L, Leu; M, Mer; N, Asn; P, Pro; Q, Gln; R, Arg; S, Ser; T, Thr; V, Vei; W, Tro: and Y, Tyr.
- 30. A W303 1A cherivativa, SY2825 (MATa ura3-1 lau2-3, 112 trp1-1 ads2-1 can1-100 sst1 & mis2A::RUS1-tsc2 his36:FUS1-HIS3), was the parent strain for the mutant search. SY2625 deviatives for the mating asseys, secreted pheromone at seys, and the pulse-charge emerments included the following streets: Y49 (stazz-1), Y115 (mitr12-1282), Y122 (sut1:14R43), Y173 (sut12-14R43), Y221 (sut12-14R43), Y220 (sut12-14R43), Y221 (sut12-14R43), Y231 (sut12-14R42), Y233 (stazz), MATe derivatives of Y233 (stazz), MATe derivatives of SY2625 included the following strains: Y199 (SY2625 made MATa), Y278 (sta22-1), Y195 (mts14::LEUZ), Y196 (std14::LEUZ), and Y197 (auti::URAS). The EG123 (MATe leu2 ura3 trp1 cant his4) genetic background was used to create a set of rasel garante bacupround was used to create a set of strains for analysis of but atta selection. EG123 de-rivatives included the following strains: Y175 (aut1 Att. EU2), Y223 (aut1 turk A), Y234 (staz3At-LEU2), and Y272 (aut1 Att. EU2 staz3At-LEU2). MATa derivatives of EG123 included the tollowing strains: Y214 (EG123 made MATa) and Y293 (ext) A::LEU2). All strains were generated by means of standard genetic or molecular methods involving the appropriate constructs (23), in particular, the axif ste23 double mutant strains were created by crossing of the appropriate MATe ste23 and MATe add mutants, followed by sporutation of the resultant diploid and isolation of the double mutant from nonparental di-type tetrada. Gene disruptions were confirmed with either PCR or Southern (DNA) analysis. 31. p129 is a YEp352 LL E. HR. A. M. Myers, T. J. Ko-erner, A. Tzagoloff, Yessi 2, 163 (1995)) plasmid con-

taining a 5.5-ldb Sel I tragment of pAXL1, p151 was

derived from p129 by insertion of a linker at the Bgl &

site within AVC.1, which led to an in-trame insertion of the hemapphatinin (HA) epitope (DQYPYDVPDYA) (29)

between arritho acids 854 and 855 of the AXL1 prod-

uct.pC225 is a KS+'(Stratagene) plasmid containing a 0.54b Barn H-Sat I tragment from pAX.1. Subs aution mutations of the proposed active site of Aid1p. were created with the use of pC225 and site-specific mulagenesis involving appropriate synthetic oligonucescient (edf-H684, 5°-GTGCTCACAAAGCGCT-GOCCHECOGGC 3": adl-E71A, 5"-MGAATCAT-GTGCGCACAAAGGTGCGC-3"; and an1-E71D, 5"-AMGANTCATGTGATCACAAAGGTGCGC-37. The mutations were confirmed by sequence analysis. After mutagenesis, the 0.44th Barn H-Msc I tragment from the mutagenized pC225 plasmids was transferred into pAXL1 to create a set of pRS316 plasmids carrying different AXL1 aloies, p124 (ax1-HS84). carrying different A/LT enems, p124 gast-model, p130 (axt1-E71A), and p132 (axt1-E71D). Similarly, a set of HA-tagged alleies carried on YEn352 were created eller replacement of the p151 Barn HA-Mac-ated eller replacement of the p151 Barn HA-Mac-ated eller replacement. fragment, to generate p151 (auti-E71A), p162 (auti-

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Quantitative Monitoring of Gene Expression Patterns with a Complementary DNA Microarray

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A high-capacity system was developed to monitor the expression of many genes in parallel. Microarrays prepared by high-speed robotic printing of complementary DNAs on glass were used for quantitative expression measurements of the corresponding genes. Because of the small format and high density of the arrays, hybridization volumes of 2 microliters could be used that enabled detection of rare transcripts in probe mixtures derived from 2 micrograms of total cellular messenger RNA. Differential expression measurements of 45 Arabidopsis genes were made by means of simultaneous, two-color

The temporal, developmental, topographical, histological, and physiological patterns in which a gene is expressed provide clues to its biological role. The large and expanding database of complementary DNA (cDNA) sequences from many organisms (1) presents the opportunity of defining these patterns at the level of the whole genome.

For these studies, we used the small flowering plant Arabidopsis thaliana as a model organism. Avabidopsis possesses many advantages for gene expression analysis, including the fact that it has the smallest genome of any higher eukaryote examined to date (2). Forty-five cloned Arabidopsis cDNAs (Table 1), including 14 complete sequences and 31 expressed sequence tags (ESTs), were used as gene-specific targets. We obtained the ESTs by selecting cDNA clones at random from an Arabidopsis cDNA library. Sequence analysis revealed that 28 of the 31 ESTs matched sequences

in the database (Table 1). Three additional cDNAs from other organisms served as controls in the experiments.

The 48 cDNAs, averaging ~1.0 kb, were amplified with the polymerase chain reaction (PCR) and deposited into individual wells of a 96-well microtiter plate. Each sample was duplicated in two adjacent wells to allow the reproducibility of the arraying and hybridization process to be tested. Samples from the microtiter plate were printed onto glass microscope slides in an area measuring 3.5 mm by 5.5 mm with the use of a high-speed arraying machine (3). The arrays were processed by chemical and heat treatment to attach the DNA sequences to the glass surface and denature them (3). Three arrays, printed in a single lot, were used for the experiments here. A single microtiter plate of PCR products provides sufficient material to print at least 500 arrays.

Fluorescent probes were prepared from total Arabidopsis mRNA (4) by a single round of reverse transcription (5). The Arabidopsis mRNA was supplemented with human acetylcholine receptor (AChR) mRNA at a dilution of 1:10,000 (w/w) before cDNA synthesis, to provide an internal standard for calibration (5). The resulting fluorescently labeled cDNA mixture was hybridized to an array at high stringency (6) and scanned

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with a laser (3). A high-sensitivity scan gave signals that saturated the detector at nearly all of the Arabidosis target sites (Fig. 1A). Calibration relative to the AChR mRNA standard (Fig. 1A) established a sensitivity limit of ~1:50,000. No detectable hybridization was observed to either the rat glucocorticoid receptor (Fig. 1A) or the yeast TRP4 (Fig. 1A) targets even at the highest scanning sensitivity. A moderate-sensitivity scan

of the same array allowed linear detection of the more abundant transcripts (Fig. 1B). Quantitation of both scans revealed a range of expression levels spanning three orders of magnitude for the 45 genes tested (Table 2). RNA blots (7) for several genes (Fig. 2) comborated the expression levels measured with the microarray to within a factor of 5 (Table 2).

Differential gene expression was investi-

gated with a simultaneous, two-color hybridization scheme, which served to minimize experimental variation inherent in the comparison of independent hybridizations. Fluorescent probes were prepared from two mRNA sources with the use of reverse transcriptase in the presence of fluorescein- and lissamine-labeled nucleotide analogs, respectively (5). The two probes were then mixed together in equal proportions, hybridized to a single array, and scanned separately for fluorescein and lissamine emission after independent excitation of the two fluorophores (3).

To test whether overexpression of a single gene could be detected in a pool of total Arabidopsu mRNA, we used a microarray to analyze a transgenic line overexpressing the single transcription factor HAT4 (8). Fluorescent probes representing mRNA from wild-type and HAT4-transgenic plants were labeled with fluorescein and lissamine, respectively; the two probes were then mixed and hybridized to a single array. An intense hybridization signal was observed at the position of the HAT4 cDNA in the lissamine-specific scan (Fig. 1D), but not in the fluorescein-specific scan of the same array (Fig. 1C). Calibration with AChR mRNA added to the fluorescein and lissamine cDNA synthesis reactions at dilutions of 1:10,000 (Fig. 1C) and 1:100 (Fig. 1D), respectively, revealed a 50-fold elevation of HAT4 mRNA in the transgenic line relative to its abundance in wild-type plants. (Table 2). This magnitude of HAT4 overexpression matched that inferred from the Northern (RNA) analysis within a factor of 2 (Fig. 2 and Table 2). Expression of all the other genes monitored on the array differed by less than a factor of 5 between HAT4transgenic and wild-type plants (Fig 1; C

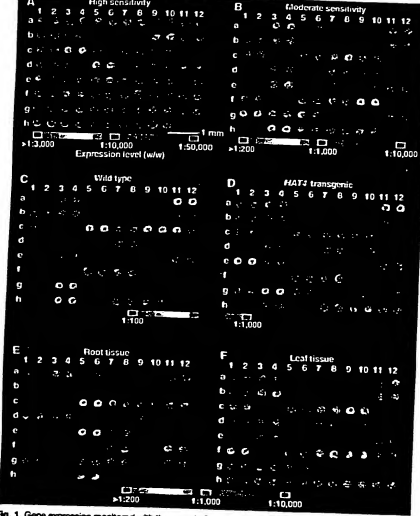


Fig. 1. Gene expression monitored with the use of cDNA microarrays. Fluorescent scans represented in pseudocolor correspond to hybridization intensities. Color bars were calibrated from the signal obtained with the use of known concentrations of human AChR mRNA in independent experiments. Numbers and letters on the axes mark the position of each cDNA. (A) High-sensitivity fluorescein can after hybridization with fluorescein-labeled cDNA derived from wild-type plants. (B) Same array as in (A) but scanned at moderate sensitivity. (C and D) A single array was probed with a 1:1 mixture of fluorescein-labeled cDNA from haT4-transgenic plants. The single array was then scanned successively to detect the fluorescein fluorescence corresponding to mRNA from wild-type plants (C) and the lissamine fluorescence corresponding to mRNA from HAT4-transgenic plants (D). (E and F) A single array was probed with a 1:1 mixture of fluorescein-labeled cDNA from root tissue and lissamine-labeled cDNA from leaf tissue. The single array was then scanned successively to detect the fluorescein fluorescence corresponding to mRNAs expressed in roots (E) and the lissamine fluorescence corresponding to mRNAs expressed in leaves (F).

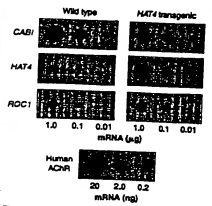


Fig. 2. Gene expression monitored with RNA (Northern) blot analysis. Designated amounts of mRNA from wild-type and HAT4-transgenic plants were spotted onto nylon membranes and probed with the cDNAs indicated. Purified human AChR mRNA was used for calibration.

and D, and Table 2). Hybridization of fluorescein-labeled glucocorticoid receptor cDNA (Fig. 1C) and lissamine-labeled TRP4 cDNA (Fig. 1D) verified the presence of the negative control targets and the lack of optical cross talk between the two fluorophores.

To explore a more complex alteration in expression patterns, we performed a second two-color hybridization experiment with fluorescein- and lissamine-labeled probes prepared from root and leaf mRNA, respectively. The scanning sensitivities for the two fluorophores were normalized by matching the signals resulting from AChR

mRNA, which was added to both cDNA synthesis reactions at a dilution of 1:1000 (Fig. 1, E and F). A comparison of the scans revealed widespread differences in gene expression between root and leaf tissue (Fig. 1, E and F). The mRNA from the light-regulated CABI gene was ~500-fold more abundant in leaf (Fig. 1F) than in root tissue (Fig. 1E). The expression of 26 other genes differed between root and leaf tissue by more than a factor of 5 (Fig. 1, E and F).

The HAT4-transgenic line we examined has elongated hypocotyls, early flowering, poor germination, and altered pigmentation (8). Although changes in expression were

observed for HAT4, large changes in expression were not observed for any of the other 44 genes we examined. This was somewhat surprising, particularly because comparative analysis of leaf and root tissue identified 27 differentially expressed genes. Analysis of an expanded set of genes may be required to identify genes whose expression changes upon HAT4 overexpression; alternatively, a comparison of mRNA populations from specific tissues of wild-type and HAT4-transgenic plants may allow identification of downstream genes.

At the current density of robotic printing, it is feasible to scale up the fabrication process to produce arrays containing 20,000 cDNA targets. At this density, a single array would be sufficient to provide gene-specific targets encompassing nearly the entire repertoire of expressed genes in the Arabidopsis genome (2). The availability of 20,274 ESTs from Arabidopsis (1, 9) would provide a rich source of templates for such studies.

The estimated 100,000 genes in the human genome (10) exceeds the number of Arabidopsis genes by a factor of 5 (2). This modest increase in complexity suggests that similar cDNA microarrays, prepared from the rapidly growing repertoire of human ESTs (1), could be used to determine the expression patterns of tens of thousands of human genes in diverse cell types. Coupling an amplification strategy to the reverse transcription reaction (11) could make it feasible to monitor expression even in minute tissue samples. A wide variety of acute and chronic physiological and pathological conditions might lead to characteristic changes in the patterns of gene expression in peripheral blood cells or other easily sampled tissues. In concert with cDNA microarrays for monitoring complex expression patterns, these tissues might therefore serve as sensitive in vivo sensors for clinical diagnosis. Microarrays of cDNAs could thus provide a useful link between human gene sequences and clinical medicine.

Table 1. Sequences contained on the cDNA microarray. Shown is the position, the known or putative function, and the accession number of each cDNA in the microarray (Fig. 1). All but three of the ESTs used in this study matched a sequence in the database. NADH, reduced form of nicotinamide adenine dinucleotide; ATPase, adenosine triphosphatase; GTP, guanosine triphosphata.

Position	AChR	Function	Accession	
81, 2 83, 4		Human AChR		
a5, 4	EST3	Actin	•	
87. 8	EST6	NADH dehydrogenase	H36236	
89, 10	AAC1	Actin 1	Z 27010	
811, 12	EST12	Unknown	M20016	
b1, 2	EST13	Actin	U36594†	
b3, 4	CABI	Chlorophyll a/o binding	T45783	
აა, 4 ა5, 6	EST17	Phosphoglycerate kinase	M85150	
~. 0 07, 8	G44	Gioberellic acid biosynthesis	T44490	
9, 10	EST19	Unknown	L37126	
11, 12	GBF-1	G-box binding factor 1	U36595†	
1, 2	EST23	Bongation factor	X63894	
	EST29	Aldolase	X52256	
3, 4	G8F-2	G-box binding factor 2	T04477	
5, 6	EST34	Chloroplast protease	X63895	
7, 8	EST35	Unknown	R87034	
. 10	EST41	Catalase	T14152	
1, 12	rGR	Rat on conscious	T22720	
. 2	EST42	Plat glucocorticoid receptor Unknown	M14053	
, 4	EST45	ATPase	U36596t	
, 6	HATT		J04185	
, 8	EST46	Homeobox-leucine zipper 1	U09332	
, 10	EST49	Light harvesting complex	T04063	
1, 12	HAT2	Unknown	T76267	
2	HAT4	Homeobox-leucine zipper 2	U09335	
4	EST50	Homeobox-leucine zipper 4	M90394	
6	HATS	PTIQSDNONDUIDIdnaga	T04344	
8	EST51	Hameabax-leucine zipper 5	M90416	
10	HAT22	UTIKNOWN	Z33675	
, 12	EST52	Homeobox-leucine zipper 22		
2	EST59	Uzygen evolving	U09336	
,		Unknown	T21749	
; ;	KNAT1	Knatted-like homeobox 1	Z34607	
3	EST60	HUBISCO SMAIL SUIDU INIT	U14174	
0	EST69	Translation elongation factor	X14564	
12	PPH1	Protein phosphatase 1	T42799	
2 2	EST70	Unknown	U34803	
4	EST75	Chloroplast protease	T44621	
5	EST78	Unknown	T43698	
	ROC1	Cyclophilin	R65481	
3	EST82	GTP binding	L14844	
10	EST83	Unknown	X59152	
12	EST84	Unknown	Z33795	
	EST91	Unknown	T45278	
ļ	EST96	Unknown	T13832	
i	SARI	Synaptobrevin	R64816	
_	EST100	Light honoration	M90418	
0	EST103	Light harvesting complex	Z18205	
12	TRP4	Light harvesting complex	X03909	
	of Stratagene (La Jolia (Yeast tryptophan biosynthesis	X04273	

*Proprietary sequence of Stratagene (La Jolla, California). The match in the database: novel EST.

Table 2. Gene expression monitoring by microarray and RNA blot analyses; tg. FA74-transgenic. See Table 1 for additional gene information. Expression levels (w/w) were calibrated with the use of known amounts of human ACNR mRNA. Values for the microarray were determined from microarray scans (Fig. 1); values for the RNA blot were determined from RNA blots (Fig. 2).

Gene	Expression level (w/w)		
	Microarray	RNA blo	
CABI CABI (1g) HAT4 HAT4 (1g) ROC1 ROC1 (1g)	1:48 1:120 1:8300 1:150 1:1200 1:260	1:83 1:150 1:6300 1:210 1:1800 1:1300	

REFERENCES AND NOTES

- 1. The current EST database (dbEST release 001495) from the National Center for Biotechnology Informs tion (Sethesda, MO) contains a total of 322,225 entries, including 255,645 from the human genome and 21,044 from Arabidopals. Access is available via he World Wide Web (http://www.ncbi.nim.nin.gov).
- 2. E. M. Mayerowitz and R. E. Pruitt. Science 229, 1214 (1985); R. E. Prutt and E. M. Mayerowitz, J. Mol. Biol. 187, 169 (1986); L Hwang et al., Plant J. 1, 357 (1991); P. Janks et al., Plant Mol. Biol. 24, 685 (1994); L. Le Guen et al., Mol. Gen. Genet. 245, 390 (1994).
- 3. D. Shalon, thesis, Stanford University (1995); and P. O. Brown, in preparation. Mic tabricated on poly-L-lyaine-costed microscope sides (Sigme) with a custom-built erraying mechine fitted with one printing tip. The tip loaded 1 µl of PCR product (0.5 mg/m) from 96-well microtter plates and deposited ~0.005 µl per side on 40 sides at a specing of 500 jum. The printed slides were rehydrated for 2 hours in a humid chember, anap-dried at 100°C for 1 min, rinsed in 0.1% SDS, and tre with 0.05% auctinic anhydride prepared in buller consisting of 50% 1-mathyl-2-pyrrolidinons and 50% boric acid. The cDNA on the sides was denstured in distilled water for 2 min at 90°C immediately before use. Microerrays were scanned with a laser cent scanner that contained a computer-controlled XY stage and a microscope objective. A mixed gas, multiline leser allowed sequential excitation of the two fluorophores. Emitted fight was split accordingth and detected with two photomultiplier tubes. Signals were read into a PC with the use of a 12-bit analog-to-digital board. Additional details of microarray tabrication and use may be obtained by means of e-mail (pbrown@crngm. stanford.edu).
- F. M. Ausubel et al., Eds., Current Protocols in Mo-lecular Biology (Greene & Wiley Interacience, New York, 1994), pp. 4.3.1-4.3.4,
- Polyaconyleted (poly4)*] mRNA was prepared from total RNA with the use of Oligotex-off resin (Clagery). Reverse transcription (RT) reactions were certed out with a StratsScript RT-PCR kit (Stratsgere) modified as follows: 50-µl reactions contained 0.1 µg/µl of Arabidopsis mRNA, 0.1 ng/µl of human AChR mRNA, 0.05 $\mu p/\mu l$ of digo(oT) (21-mer), 1× first strand buffer, 0.03 U/ μl of ribonuclease block, 500 what come, was our or monuclesse block, 500 μM decoyguanceine triphosphate, 500 μM decoyguanceine triphosphate, 500 μM duranceine triphosphate (dCTP), 40 μM θμοσιασοίη-12-dCTP (or lissamine-5-dCTP), and 0.03 U/µl of StrateScript reverse transcriptase. Re re incubated for 60 min at 37°C, precipitated with ethanol, and resuspended in 10 µl of TE (10 mM tris-HCl and 1 mM EDTA, pH 8.0). Samples were than heated for 3 min at 94°C and chilled on ice. The RNA was degraded by adding 0.25 pl of 10 N NaOH followed by a 10-min incubation at 37° are neutralized by addition of 2.5 μ l of 1 M tris-CI (pH 8.0) and 0.25 µJ of 10 N HCI and pracioitsted with etherol. Pelets were washed with 70% ethanol, dried to completion in a speedvac. pended in 10 µl of H₂O, and reduced to 3.0 µl in a speedvec. Puorescent nucleotide analogs were obtained from New England Nuclear (DuPont).
- 6. Hybridization reactions contained 1.0 µl of fluorescent cONA synthesis product (5) and 1.0 µJ of hybridization buffer (10× saline sodium citrate (SSC) and 0.2% SDS). The 2.0-µi probe mixtures were aliquoted onto the microarray surface and covered with cover slips (12 mm round). Arrays were transferred to a hybrid-ization chamber (3) and incubated for 18 hours at 65°C. Arrays were washed for 5 min at room tempersture (25°C) in low-stringency wash buffer (1 × SSC and 0.1% SDS), then for 10 min at room temperature in high-stringency wash butter (0.1% SSC and 0.1% SDS). Arrays were scanned in 0.1 × SSC with the use scence leser-scenning device (3).
- 7. Samples of poly(A)* mRNA (4, 5) were spotted onto nylon membranes (Nyltran) and crosslinked with ultraviolet light with the use of a Stratalinker 1800 (Stratagene). Probes were prepared by random priming with the use of a Prime-It is kit (Stratagene) in the presence of PPPJCATP. Hybridizations were carried out according to the instructions of the manu-

- tecturer. Quantitation was performed on a Phosphorimager (Molecular Dynamics).
- 8. M. Schene and R. W. Davis, Proc. Natl. Acad. Sci. U.S.A. 89, 3894 (1992); M. Schena, A. M. Lloyd, R. W. Davis, Genes Dev. 7, 367 (1993); M. Schena and R. W. Davis, Proc. Natl. Acad. Sci. U.S.A. 91, 8393
- 9. H. Hofte et al., Plant J. 4, 1051 (1993); T. Newman et al., Plant Physiol. 106, 1241 (1994).
- 10. N. E. Morton, Proc. Natl. Acad. Sci. U.S.A. 88, 7474 (1991); E. D. Green and R. H. Waterston, J. Am. Med. Assoc. 266, 1966 (1991); C. Bellenne-Chantalot, Cell 70, 1059 (1992); D. R. Cox et al., Science 265, 2031 (1994).
- Hi et al., Proc. Netl. Aced. Sci. U.S.A. 85, 5698 (1988).
- 12. The laser \$uorescent scenner was de d in collaboration with S. Smith of Stanford Unive ally. Scanner and analysis activers was developed by R. X. Xia. The aucoinic arthydride reaction was augo ed by J. Muligan and J. Van Ness of Darwin Molecular Corporation, Thereis to S. Theologie, C. Somerve Yarramoto, and members of the laboratories of R.W.D. and P.O.B. for critical comments. Supported by the Howard Hughes Medical institute and by grants from MRH [R21HG00450] P.O.B.) and R37AG00198 (R.W.D.)] and from NSF (MCB9106011) (R.W.D.) and by an NSF graduate fellowship (D.S.). P.O.B. is an art investigator of the Howard Hughes Medical
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Gene Therapy in Peripheral Blood Lymphocytes and Bone Marrow for ADA Immunodeficient Patients

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Adenosine dearninase (ADA) deficiency results in severe combined immunodeficiency, the first genetic disorder treated by gene therapy. Two different retroviral vectors were used to transfer ex vivo the human ADA minigene into bone marrow cells and peripheral blood lymphocytes from two patients undergoing exogenous enzyme replacement therapy. After 2 years of treatment, long-term survival of T and B lymphocytes, marrow cells, and granulocytes expressing the transferred ADA gene was demonstrated and resulted in normalization of the immune repertoire and restoration of cellular and humoral immunity. After discontinuation of treatment, T lymphocytes, derived from transduced peripheral blood lymphocytes, were progressively replaced by marrow-derived T cells in both patients. These results indicate successful gene transfer into long-lasting progenitor cells, producing a functional multilineage progeny.

Severe combined immunodeficiency associated with inherited deficiency of ADA (1) is usually fatal unless affected children are kept in protective isolation or the immune system is reconstituted by bone marrow transplantation from a human leukocyte antigen (HLA)-identical sibling donor (2). This is the therapy of choice, although it is available only for a minority of patients. In recent years, other forms of therapy have been developed, including transplants from haploidentical donors (3, 4), exogenous ensyme replacement (5), and somatic-cell gene therapy (6-9).

We previously reported a preclinical model in which ADA gene transfer and expression

blood lymphocytes (PBLs) in immunodeficient mice in vivo (10, 11). On the basis of these preclinical results, the clinical application of gene therapy for the treatment of ADA SCID (severe combined immunodeficiency disease) patients who previously failed exogenous enzyme replacement therapy was approved by our Institutional Ethical Committees and by the Italian National Committee for Bioethics (12). In addition to evaluating the safety and efficacy of the gene therapy procedure, the aim of the study was to define the relative role of PBLs and hematopoietic stem cells in the long-term reconstitution of immune functions after retroviral vector-mediated ADA gene transfer. For this purpose, two structurally identical vectors expressing the human ADA complementary DNA (cDNA), distinguishable by the presence of alternative restriction sites in a nonfunctional region of the viral long-terminal repeat

(LTR), were used to transduce PBLs and bone marrow (BM) cells independently. This pro-

cedure allowed identification of the origin of

successfully restored immune functions in hu-

man ADA-deficient (ADA⁻) peripheral

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Differential gene expression in drug metabolism and toxicology: practicalities, problems and potential

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- 1. An important feature of the work of many molecular biologists is identifying which genes are switched on and off in a cell under different environmental conditions or subsequent to xenobiotic challenge. Such information has many uses, including the deciphering of molecular pathways and facilitating the development of new experimental and diagnostic procedures. However, the student of gene hunting should be forgiven for perhaps becoming confused by the mountain of information available as there appears to be almost as many methods of discovering differentially expressed genes as there are research groups using the technique.
- 2. The aim of this review was to clarify the main methods of differential gene expression analysis and the mechanistic principles underlying them. Also included is a discussion on some of the practical aspects of using this technique. Emphasis is placed on the so-called 'open' systems, which require no prior knowledge of the genes contained within the study model. Whilst these will eventually be replaced by 'closed' systems in the study of human, mouse and other commonly studied laboratory animals, they will remain a powerful tool for those examining less fashionable models.
- 3. The use of suppression-PCR subtractive hybridization is exemplified in the identification of up- and down-regulated genes in rat liver following exposure to phenobarbital, a well-known inducer of the drug metabolizing enzymes.
- 4. Differential gene display provides a coherent platform for building libraries and microchip arrays of 'gene fingerprints' characteristic of known enzyme inducers and xenobiotic toxicants, which may be interrogated subsequently for the identification and characterization of xenobiotics of unknown biological properties.

Introduction

It is now apparent that the development of almost all cancers and many non-neoplastic diseases are accompanied by altered gene expression in the affected cells compared to their normal state (Hunter 1991, Wynford-Thomas 1991, Vogelstein and Kinzler 1993, Semenza 1994, Cassidy 1995, Kleinjan and Van Hegningen 1998). Such changes also occur in response to external stimuli such as pathogenic microorganisms (Rohn et al. 1996, Singh et al. 1997, Griffin and Krishna 1998, Lunney 1998) and xenobiotics (Sewall et al. 1995, Dogra et al. 1998, Ramana and Kohli 1998), as well as during the development of undifferentiated cells (Hecht 1998, Rudin and Thompson 1998, Schneider-Maunoury et al. 1998). The potential medical and therapeutic benefits of understanding the molecular changes which occur in any given cell in progressing from the normal to the 'altered' state are enormous. Such profiling essentially provides a 'fingerprint' of each step of a

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cell's development or response and should help in the elucidation of specific and sensitive biomarkers representing, for example, different types of cancer or previous exposure to certain classes of chemicals that are enzyme inducers.

In drug metabolism, many of the xenobiotic-metabolizing enzymes (including the well-characterized isoforms of cytochrome P450) are inducible by drugs and chemicals in man (Pelkonen et al. 1998), predominantly involving transcriptional activation of not only the cognate cytochrome P450 genes, but additional cellular proteins which may be crucial to the phenomenon of induction. Accordingly, the development of methodology to identify and assess the full complement of genes that are either up- or down-regulated by inducers are crucial in the development of knowledge to understand the precise molecular mechanisms of enzyme induction and how this relates to drug action. Similarly, in the field of chemical-induced toxicity, it is now becoming increasingly obvious that most adverse reactions to drugs and chemicals are the result of multiple gene regulation, some of which are causal and some of which are casually-related to the toxicological phenomenon per se. This observation has led to an upsurge in interest in gene-profiling technologies which differentiate between the control and toxin-treated gene pools in target tissues and is, therefore, of value in rationalizing the molecular mechanisms of xenobioticinduced toxicity. Knowledge of toxin-dependent gene regulation in target tissues is not solely an academic pursuit as much interest has been generated in the pharmaceutical industry to harness this technology in the early identification of toxic drug candidates, thereby shortening the developmental process and contributing substantially to the safety assessment of new drugs. For example, if the gene profile in response to say a testicular toxin that has been well-characterized in vivo could be determined in the testis, then this profile would be representative of all new drug candidates which act via this specific molecular mechanism of toxicity, thereby providing a useful and coherent approach to the early detection of such toxicants. Whereas it would be informative to know the identity and functionality of all genes up/down regulated by such toxicants, this would appear a longer term goal, as the majority of human genes have not yet been sequenced, far less their functionality determined. However, the current use of gene profiling yields a pattern of gene changes for a xenobiotic of unknown toxicity which may be matched to that of wellcharacterized toxins, thus alerting the toxicologist to possible in vivo similarities between the unknown and the standard, thereby providing a platform for more extensive toxicological examination. Such approaches are beginning to gain momentum, in that several biotechnology companies are commercially producing 'gene chips' or 'gene arrays' that may be interrogated for toxicity assessment of xenobiotics. These chips consist of hundreds/thousands of genes, some of which are degenerate in the sense that not all of the genes are mechanistically-related to any one toxicological phenomenon. Whereas these chips are useful in broad-spectrum screening, they are maturing at a substantial rate, in that gene arrays are now becoming more specific, e.g. chips for the identification of changes in growth factor families that contribute to the aetiology and development of chemically-induced neoplasias.

Although documenting and explaining these genetic changes presents a formidable obstacle to understanding the different mechanisms of development and disease progression, the technology is now available to begin attempting this difficult challenge. Indeed, several 'differential expression analysis' methods have been developed which facilitate the identification of gene products that demonstrate

altered expression in cells of one population compared to another. These methods have been used to identify differential gene expression in many situations, including invading pathogenic microbes (Zhao et al. 1998), in cells responding to extracellular and intracellular microbial invasion (Duguid and Dinauer 1990, Ragno et al. 1997, Maldarelli et al. 1998), in chemically treated cells (Syed et al. 1997, Rockett et al. 1999), neoplastic cells (Liang et al. 1992, Chang and Terzaghi-Howe 1998), activated cells (Gurskaya et al. 1996, Wan et al. 1996), differentiated cells (Hara et al. 1991, Guimaraes et al. 1995a, b), and different cell types (Davis et al. 1984, Hedrick et al. 1984, Xhu et al. 1998). Although differential expression analysis technologies are applicable to a broad range of models, perhaps their most important advantage is that, in most cases, absolutely no prior knowledge of the specific genes which are up- or down-regulated is required.

The field of differential expression analysis is a large and complex one, with many techniques available to the potential user. These can be categorized into several methodological approaches, including:

(1) Differential screening,

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- (2) Subtractive hybridization (SH) (includes methods such as chemical cross-linking subtraction—CCLS, suppression-PCR subtractive hybridization—SSH, and representational difference analysis—RDA).
- (3) Differential display (DD),
- (4) Restriction endonuclease facilitated analysis (including serial analysis of gene expression—SAGE—and gene expression fingerprinting—GEF),
- (5) Gene expression arrays, and
- (6) Expressed sequence tag (EST) analysis.

The above approaches have been used successfully to isolate differentially expressed genes in different model systems. However, each method has its own subtle (and sometimes not so subtle) characteristics which incur various advantages and disadvantages. Accordingly, it is the purpose of this review to clarify the mechanistic principles underlying the main differential expression methods and to highlight some of the broader considerations and implications of this very powerful and increasingly popular technique. Specifically, we will concentrate on the so-called 'open' systems, namely those which do not require any knowledge of gene sequences and, therefore, are useful for isolating unknown genes. Two 'closed' systems (those utilising previously identified gene sequences), EST analysis and the use of DNA arrays, will also be considered briefly for completeness. Whilst emphasis will often be placed on suppression PCR subtractive hybridization (SSH, the approach employed in this laboratory), it is the aim of the authors to highlight, wherever possible, those areas of common interest to those who use, or intend to use, differential gene expression analysis.

Differential cDNA library screening (DS)

Despite the development of multiple technological advances which have recently brought the field of gene expression profiling to the forefront of molecular analysis, recognition of the importance of differential gene expression and characterization of differentially expressed genes has existed for many years. One of the original approaches used to identify such genes was described 20 years ago by St John and Davis (1979). These authors developed a method, termed 'differential plaque filter

hybridization, which was used to isolate galactose-inducible DNA sequences from yeast. The theory is simple: a genomic DNA library is prepared from normal, unstimulated cells of the test organism/tissue and multiple filter replicas are prepared! These replica blots are probed with radioactively (or otherwise) labelled complex cDNA probes prepared from the control and test cell mRNA populations. Those mRNAs which are differentially expressed in the treated cell population will show a positive signal only on the filter probed with cDNA from the treated cells. Furthermore, labelled cDNA from different test conditions can be used to probe multiple blots, thereby enabling the identification of mRNAs which are only upregulated under certain conditions. For example, St John and Davis (1979) screened replica filters with acetate-, glucose- and galactose-derived probes in order to obtain genes induced specifically by galactose metabolism. Although groundbreaking in its time this method is now considered insensitive and time-consuming, as up to 2 months are required to complete the identification of genes which are differentially expressed in the test population. In addition, there is no convenient way to check that the procedure has worked until the whole process has been completed. about of the region with the first of the section and the section and the region of

Subtractive Hybridization (SH)

The developing concept of differential gene expression and the success of early approaches such as that described by St John and Davis (1979) soon gave rise to a search for more convenient methods of analysis. One of the first to be developed was SH, numerous variations of which have since been reported (see below). In general, this approach involves hybridization of mRNA/cDNA from one population (tester) to excess mRNA/cDNA from another (driver), followed by separation of the unhybridized tester fraction (differentially expressed) from the hybridized common sequences. This step has been achieved physically, chemically and through the use of selective polymerase chain reaction (PCR) techniques.

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Physical separation

Original subtractive hybridization technology involved the physical separation of hybridized common species from unique single stranded species. Several methods of achieving this have been described, including hydroxyapatite chromatography (Sargent and Dawid 1983), avidin-biotin technology (Duguid and Dinauer 1990) and oligodT-latex separation (Hara et al. 1991). In the first approach, common mRNA species are removed by cDNA (from test cells)-mRNA (from control cells) subtractive hybridization followed by hydroxyapatite chromatography, as hydroxyapatite specifically adsorbs the cDNA-mRNA hybrids. The unabsorbed cDNA is then used either for the construction of a cDNA library of differentially expressed genes (Sargent and Dawid 1983, Schneider et al. 1988) or directly as a probe to screen a preselected library (Zimmerman et al. 1980, Davis et al. 1984, Hedrick et al. 1984). A schematic diagram of the procedure is shown in figure 1.

Less rigorous physical separation procedures coupled with sensitivity enhancing PCR steps were later developed as a means to overcome some of the problems encountered with the hydroxyapatite procedure. For example, Daguid and Dinauer (1990) described a method of subtraction utilizing biotin-affinity systems as a means to remove hybridized common sequences. In this process, both the control and tester mRNA populations are first converted to cDNA and an adaptor ('oligovector',

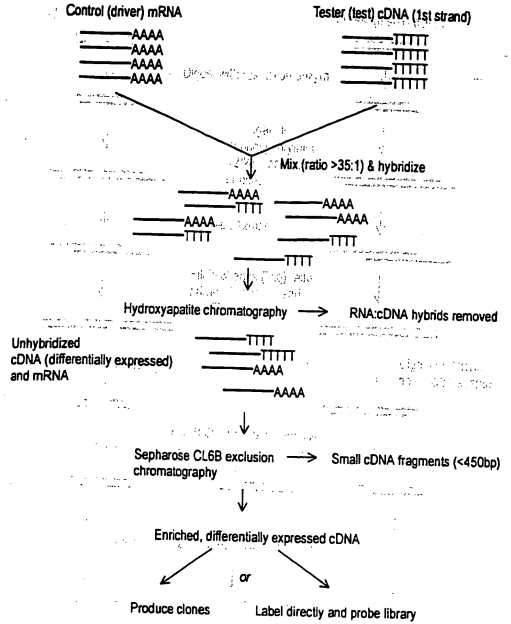


Figure 1. The hydroxyapatite method of subtractive hybridization. cDNA derived from the treated/altered (tester) population is mixed with a large excess of mRNA from the control (driver) population. Following hybridization, mRNA-cDNA hybrids are removed by hydroxyapatite chromatography. The only cDNAs which remain are those which are differentially expressed in the treated/altered population. In order to facilitate the recovery of full length clones, small cDNA fragments are removed by exclusion chromatography. The remaining cDNAs are then cloned into a vector for sequencing, or labelled and used directly to probe a library, as described by Sargent and Dawid (1983).

containing a restriction site) ligated to both sides. Both populations are then amplified by PCR, but the driver cDNA population is subsequently digested with the adaptor-containing restriction endonuclease. This serves to cleave the oligovector and reduce the amplification potential of the control population. The digested control population is then biotinylated and an excess mixed with tester cDNA. Following denaturation and hybridization, the mix is applied to a biocytin column (streptavidin may also be used) to remove the control population, including heteroduplexes formed by annealing of common sequences from the tester population. The procedure is repeated several times following the addition of fresh

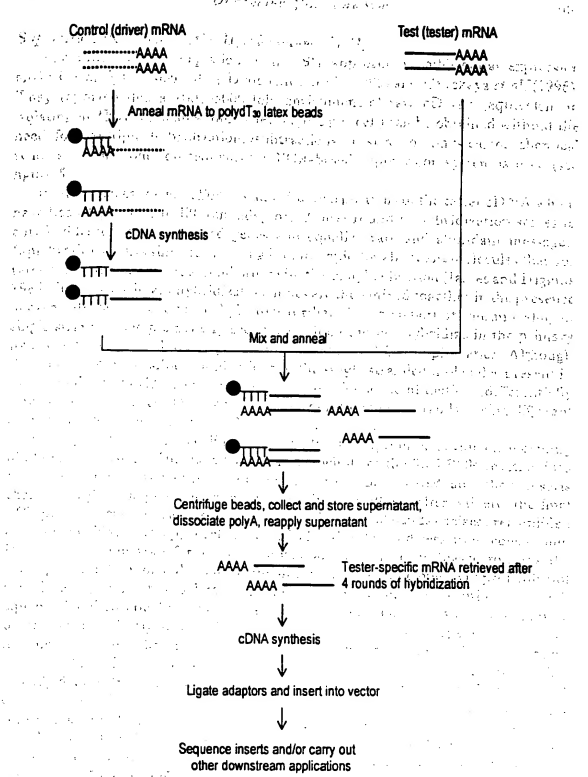


Figure 2. The use of oligodT₂₀ latex to perform subtractive hybridization. mRNA extracted from the control (driver) population is converted to anchored cDNA using polydT oligonucleotides attached to latex beads. mRNA from the treated/altered (tester) population is repeatedly hybridized against an excess of the anchored driver cDNA. The final population of mRNA is tester specific and can be converted into cDNA for cloning and other downstream applications, as described by Hara et al. (1991).

control cDNA. In order to further enrich those species differentially expressed in the tester cDNA, the subtracted tester population is amplified by PCR following every second subtraction cycle. After six cycles of subtraction (three reamplification steps) the reaction mix is ligated into a vector for further analysis.

In a slightly different approach, Hara et al. (1991) utilized a method whereby oligo(dT₃₀) primers attached to a latex substrate are used to first capture mRNA extracted from the control population. Following 1st strand cDNA synthesis, the RNA strand of the heteroduplexes is removed by heat denaturation and centrifugation (the cDNA-oligotex-dT₃₀ forms a pellet and the supernatant is removed). A quantity of tester mRNA is then repeatedly hybridized to the immobilized control (driver) cDNA (which is present in 20-fold excess). After several rounds of hybridization the only mRNA molecules left in the tester mRNA population are those which are not found in the driver cDNA-oligotex-dT₃₀ population. These tester-specific mRNA species are then converted to cDNA and, following the addition of adaptor sequences, amplified by PCR. The PCR products are then ligated into a vector for further analysis using restriction sites incorporated into the PCR primers. A schematic illustration of this subtraction process is shown in figure 2.

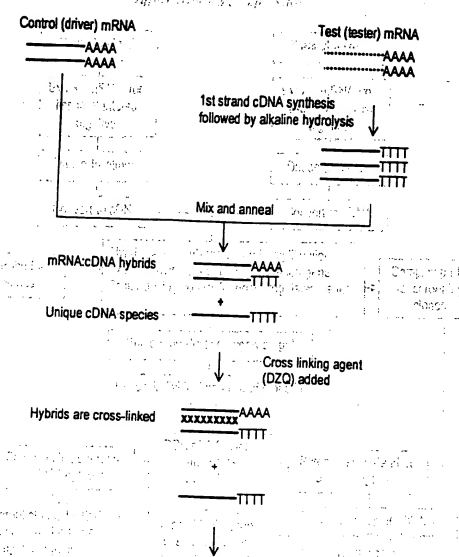
However, all these methods utilising physical separation have been described as inefficient due to the requirement for large starting amounts of mRNA, significant loss of material during the separation process and a need for several rounds of hybridization. Hence, new methods of differential expression analysis have recently been designed to eliminate these problems.

Chemical Cross-Linking Subtraction (CCLS)

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In this technique, originally described by Hampson et al. (1992), driver mRNA is mixed with tester cDNA (1st strand only) in a ratio of > 20:1. The common sequences form cDNA:mRNA hybrids, leaving the tester specific species as single stranded cDNA. Instead of physically separating these hybrids, they are inactivated chemically using 2,5 diaziridinyl-1,4-benzoquinone (DZQ). Labelled probes are then synthesized from the remaining single stranded cDNA species (unreacted mRNA species remaining from the driver are not converted into probe material due to specificity of Sequenase T7 DNA polymerase used to make the probe) and used to screen a cDNA library made from the tester cell population. A schematic diagram of the system is shown in figure 3.

It has been shown that the differentially expressed sequences can be enriched at least 300-fold with one round of subtraction (Hampson et al. 1992), and that the technique should allow isolation of cDNAs derived from transcripts that are present at less than 50 copies per cell. This equates to genes at the low end of intermediate abundance (see table 1). The main advantages of the CCLS approach are that it is rapid, technically simple and also produces fewer false positives than other differential expression analysis methods. However, like the physical separation protocols, a major drawback with CCLS is the large amount of starting material required (at least 10 µg RNA). Consequently, the technique has recently been refined so that a renewable source of RNA can be generated. The degenerate random oligonucleotide primed (DROP) adaptation (Hampson et al. 1996, Hampson and Hampson 1997) uses random hexanucleotide sequences to prime solid phase-synthesized cDNA. Since each primer includes a T7 polymerase promotor sequence



Probes synthesised from single stranded cDNA species and used to probe cDNA library

Figure 3. Chemical cross-linking subtraction. Excess driver mRNA is mixed with 1st strand tester cDNA. The common sequences form mRNA:cDNA hybrids which are cross linked with 2,5 diaziridinyl-1,4-benzoquinone (DZQ) and the remaining cDNA sequences are differentially expressed in the tester population. Probes are made from these sequences using Sequenase 2.0 DNA polymerase, which lacks reverse transcriptase activity and, therefore, does not react with the remaining mRNA molecules from the driver. The labelled probes are then used to screen a cDNA library for clones of differentially expressed sequences. Adapted from Walter et al. (1996), with

Table 1. The abundance of mRNA species and classes in a typical mammalian cell.

mRNA class	Copies of each species/cell	No. of mRNA species in class	Mean % of each species in class	Mean mass (ng) of each species/µg total RNA
Abundant	12 000	4	3.3	1.65
Intermediate	300	500	0.08	0.04
Rare	15	11000	0.004	0.002

Modified from Bertioli et al. (1995).

at the 5'end, the final pool of random cDNA fragments is a PCR-renewable cDNA population which is representative of the expressed gene pool and can be used to synthesize sense RNA for use as driver material. Furthermore, if the final pool of random cDNA fragments is reamplified using biotinylated T7 primer and random hexamer, the product can be captured with streptavidin beads and the antisense strand eluted for use as tester. Since both target and driver can be generated from the same DROP product, subtraction can be performed in both directions (i.e. for up- and down-regulated species) between two different DROP products.

Representational Difference Analysis (RDA)

RDA of cDNA (Hubank and Schatz 1994) is an extension of the technique originally applied to genomic DNA as a means of identifying differences between two complex genomes (Lisitsyn et al. 1993). It is a process of subtraction and amplification involving subtractive hybridization of the tester in the presence of excess driver. Sequences in the tester that have homologues in the driver are rendered unamplifiable, whereas those genes expressed only in the tester retain the ability to be amplified by PCR. The procedure is shown schematically in figure 4.

In essence, the driver and tester mRNA populations are first converted to cDNA and amplified by PCR following the ligation of an adaptor. The adaptors are then removed from both populations and a new (different) adaptor ligated to the amplified tester population only. Driver and tester populations are next melted and hybridized together in a ratio of 100:1. Following hybridization, only tester: tester homohybrids have 5'adaptors at each end of the DNA duplex and can, thus, be filled in at both 3' ends. Hence, only these molecules are amplified exponentially during the subsequent PCR step. Although tester: driver heterohybrids are present, they only amplify in a linear fashion, since the strand derived from the driver has no adaptor to which the primer can bind. Driver: driver heterohybrids have no adaptors and, therefore, are not amplified. Single stranded molecules are digested with mung bean nuclease before a further PCR-enrichment of the tester:tester homohybrids. The adaptors on the amplified tester population are then replaced and the whole process repeated a further two or three times using an increasing excess of driver (Hubank and Shatz used a tester:driver ratio of 1:400, 1:80000 and 1:800000 for the second, third and fourth hybridizations, respectively). Different adaptors are ligated to the tester between successive rounds of hybridization and amplification to prevent the accumulation of PCR products that might interfere with subsequent amplifications. The final display is a series of differentially expressed gene products easily observable on an ethidium bromide gel.

The main advantages of RDA are that it offers a reproducible and sensitive approach to the analysis of differentially expressed genes. Hubank and Schatz (1994) reported that they were able to isolate genes that were differentially expressed in substantially less than 1% of the cells from which the tester is derived. Perhaps the main drawback is that multiple rounds of ligation, hybridization, amplifiation and digestion are required. The procedure is, therefore, lengthier than many other differential display approaches and provides more opportunity for operator-induced error to occur. Although the generation of false positives has been noted, this has been solved to some degree by O'Neill and Sinclair (1997) through the use of HPLC-purified adaptors. These are free of the truncated adaptors which appear to be a major source of the false positive bands. A very similar technique to RDA, termed linker capture subtraction (LCS) was described by Yang and Sytowski (1996).

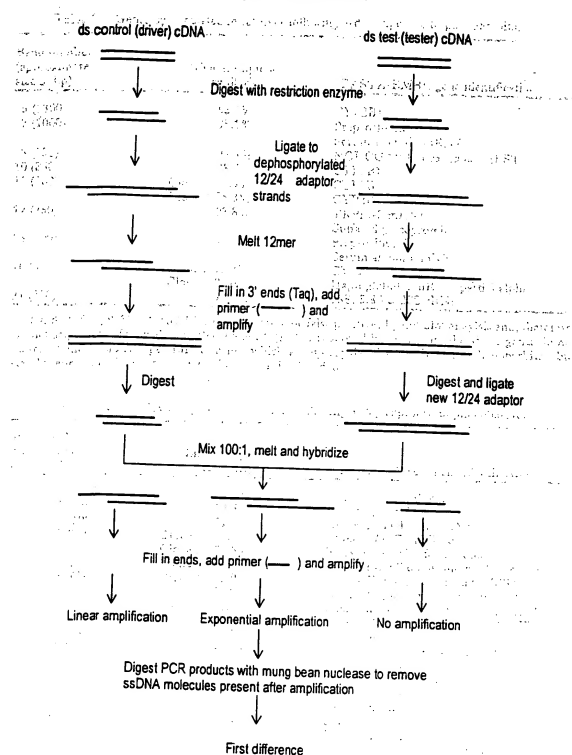


Figure 4. The representational difference analysis (RDA) technique. Driver and tester cDNA are digested with a 4-cutter restriction enzyme such as DpnII. The 1st set of 12/24 adaptor strands (oligonucleotides) are ligated to each other and the digested cDNA products. The 12mer is subsequently melted away and the 3'ends filled in using Taq DNA polymerase. Each cDNA population is then amplified using PCR, following which the 1st set of adaptors is removed with DpnII. A second set of 12/24 adaptor strands is then added to the amplified tester cDNA population, after which the tester is hybridized against a large excess of driver. The 12mer adaptors are melted and the 3' ends filled in as before. PCR is carried out with primers identical to the new 24mer adaptor. Thus, the only hybridization products which are exponentially amplified are those which are tester: tester combinations. Following PCR, ssDNA products are removed with mung bean nuclease, leaving the 'first difference product'. This is digested and a third set of 12/24 adaptors added before repeating the subtraction process from the hybridization stage. The process is repeated to the 3rd or 4th difference product, as described by Lisitsyn et al.

Suppression PCR Subtractive Hybridization (SSH)

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The most recent adaptation of the SH approach to differential expression analysis was first described by Diatchenko et al. (1996) and Gurskaya et al. (1996). They reported that a 1000-5000 fold enrichment of rare cDNAs (equivalent to isolating mRNAs present at only a few copies per cell) can be obtained without the need for multiple hybridizations/subtractions. Instead of physical or chemical removal of the common sequences, a PCR-based suppression system is used (see figure 5).

In SSH, excess driver cDNA is added to two portions of the tester cDNA which have been ligated with different adaptors. A first round of hybridization serves to enrich differentially expressed genes and equalize fare and abundant messages. Equalization occurs since reannealing is more rapid for abundant molecules than for rarer molecules due to the second order kinetics of hybridization (James and Higgins 1985). The two primary hybridization mixes are then mixed together in the presence of excess driver and allowed to hybridize further. This step permits the annealing of single stranded complementary sequences which did not hybridize in the primary hybridization, and in doing so generates templates for PCR amplification. Although there are several possible combinations of the single stranded molecules present in the secondary hybridization mix, only one particular combination (differentially expressed in the tester cDNA composed of complimentary strands having different adaptors) can amplify exponentially.

Having obtained the final differential display, two options are available if cloning of cDNAs is desired. One is to transform the whole of the final PCR reaction into competent cells. Transformed colonies can then be isolated and their inserts characterized by sequencing, restriction analysis or PCR. Alternatively, the final PCR products can be resolved on a gel and the individual bands excised, reamplified and cloned. The first approach is technically simpler and less time consuming. However, ligation/transformation reactions are known to be biased towards the cloning of smaller molecules, and so the final population of clones will probably not contain a representative selection of the larger products. In addition, although equalization theoretically occurs, observations in this laboratory suggest that this is by no means perfectly accomplished. Consequently, some gene species are present in a higher number than others and this will be represented in the final population of clones. Thus, in order to obtain a substantial proportion of those gene species that actually demonstrate differential expression in the tester population, the number of clones that will have to be screened after this step may be substantial. The second approach is initially more time consuming and technically demanding. However, it would appear to offer better prospects for cloning larger and low abundance gel products. In addition, one can incorporate a screening step that differentiates different products of different sequences but of the same size (HA-staining, see later). In this way, a good idea of the final number of clones to be isolated and identified can be achieved.

An alternative (or even complementary) approach is to use the final differential display reaction to screen a cDNA library to isolate full length clones for further characterization, or a DNA array (see later) to quickly identify known genes. SSH has been used in this laboratory to begin characterization of the short-term gene expression profiles of enzyme-inducers such as phenobarbital (Rockett et al. 1997) and Wy-14,643 (Rockett et al. unpublished observations). The isolation of differentially expressed genes in this manner enables the construction of a fingerprint

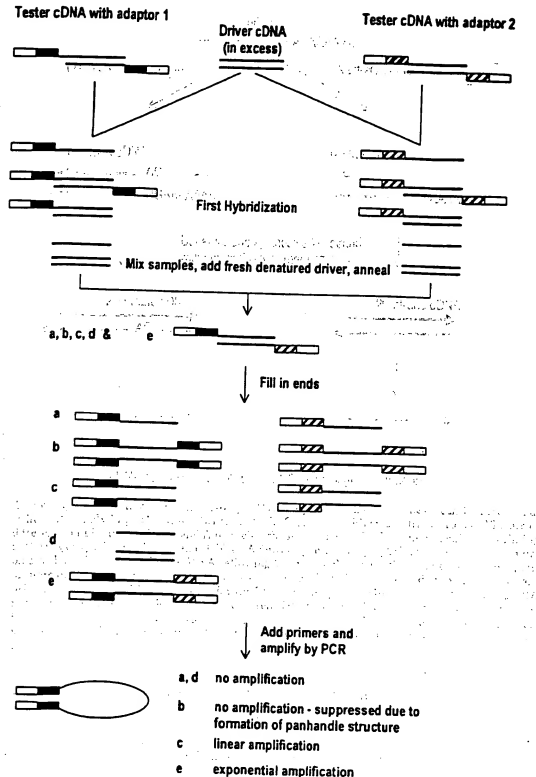


Figure 5. PCR-select cDNA subtraction. In the primary hybridization, an excess of driver cDNA is added to each tester cDNA population. The samples are heat denatured and allowed to hybridize for between 3 and 8 h. This serves two purposes: (1) to equalize rare and abundant molecules; and (2) to enrich for differentially expressed sequences—cDNAs that are not differentially expressed form type c molecules with the driver. In the secondary hybridization, the two primary hybridizations are mixed together without denaturing. Fresh denatured driver can also be added at this point to allow further enrichment of differentially expressed sequences. Type e molecules are formed in this secondary hybridization which are subsequently amplified using two rounds of PCR. The final products can be visualized on an agarose gel, labelled directly or cloned into a vector for downstream manipulation. As described by Diatchenko et al. (1996) and Gurskaya et al. (1996), with permission.

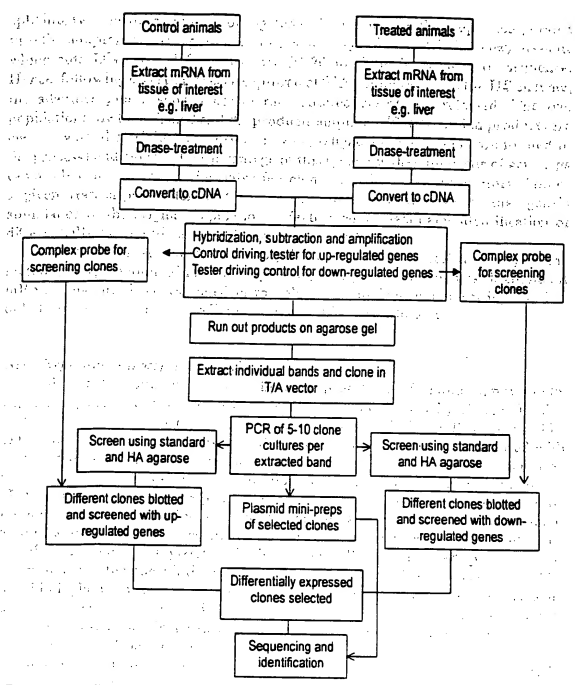


Figure 6. Flow diagram showing method used in this laboratory to isolate and identify clones of genes which are differentially expressed in rat liver following short term exposure to the enzyme inducers, phenobarbital and Wy-14,643.

of expressed genes which are unique to each compound and time/dose point. Such information could be useful in short-term characterization of the toxic potential of new compounds by comparing the gene-expression profiles they elicit with those produced by known inducers. Figure 6 shows a flow diagram of the method used to isolate, verify and clone differentially expressed genes, and figure 7 shows expression profiles obtained from a typical SSH experiment. Subsequent sub-cloning of the individual bands, sequencing and gene data base interrogation reveals many genes which are either up- or down-regulated by phenobarbital in the rat (tables 2 and 3).

One of the advantages in using the SSH approach is that no prior knowledge is required of which specific genes are up/down-regulated subsequent to xenobiotic

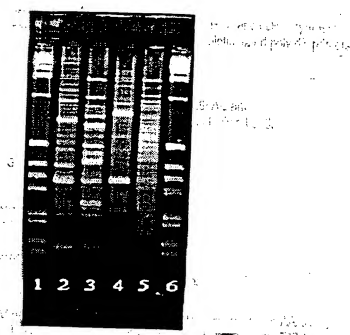


Figure 7. SSH display patterns obtained from rat liver following 3-day treatment with WY-14,643 or phenobarbital. mRNA extracted from control and treated livers was used to generate the differential displays using the PCR-Select cDNA subtraction kit (Clontech). Lane: 1—1kb ladder; 2—genes upregulated following Wy,14—643 treatment; 3—genes downregulated following Wy,14—643 treatment; 4—genes upregulated following phenobarbital treatment; 5—genes downregulated following phenobarbital treatment; 5—genes al. (1997), with permission.

exposure, and an almost complete complement of genes are obtained. For example, the peroxisome proliferator and non-genotoxic hepatocarcinogen Wy,14,643, upregulates at least 28 genes and down-regulates at least 15 in the rat (a sensitive species) and produces 48 up- and 37 down-regulated genes in the guinea pig, a resistant species (Rockett, Swales, Esda and Gibson, unpublished observations). One of these genes, CD81, was up-regulated in the rat and down-regulated in the guinea pig following Wy-14,643 treatment. CD81 (alternatively named TAPA-1) is a widely expressed cell surface protein which is involved in a large number of cellular processes including adhesion, activation, proliferation and differentiation (Levy et al. 1998). Since all of these functions are altered to some extent in the phenomena of hepatomegaly and non-genotoxic hepatocarcinogenesis, it is intriguing, and probably mechanistically-relevant, that CD81 expression is differentially regulated in a resistant and susceptible species. However, the down-side of this approach is that the majority of genes can be sequenced and matched to database sequences, but the latter are predominantly expressed sequence tags or genes of completely unknown function, thus partially obscuring a realistic overall assessment of the critical genes of genuine biological interest. Notwithstanding the lack of complete funtional identification of altered gene expression, such gene profiling studies essentially provides a 'molecular fingerprint' in response to xenobiotic challenge, thereby serving as a mechanistically-relevant platform for further detailed investigations.

Differential Display (DD)

Originally described as 'RNA fingerprinting by arbitrarily primed PCR' (Liang and Pardee 1992) this method is now more commonly referred to as 'differential

Table 2. Genes up-regulated in rat liver following 3-day exposure to phenobarbital.

		in the second of
5 (1300)	93.5%	CYP2B1 (F 1 - FO F Bruns C ORD) o
O But something		Preproalbumin Serum albumin mRNA NCI-CGAP-Pr1 H. sapiens (EST) CYP2B1 CYP2B1 CYP2B2
8 (950)	98.3%	NCI-CGAP-Pr1 H. socient (FST)
10 (850)	95.7%	CYP2B1
11 (800)	Clone 1 94.9%	the teCYP2B1s of the end of the second of th
20 mm	Clone 2 75.3%	CYP2B2
12 (750)	93.8%	CYP2B1 CYP2B2 TRPM-2 mRNA
d'e venous test and the		TRPM-2 mRNA Sulfated glycoprotein CS
13 (000)	92.9%	Sulfated glycoprotein CCp. 18 (19) 18
16 /FEN	61-10-00	Serum albumin mRNA
10 (33)	Clone 1 95.2%	CYP2B1
24/250) (2.4) (2.4)	Cione 2 93.6%	Serum albumin mRNA CYP2B1 Haptoglobulin mRNA partial alpha 18S, 5.8S & 28S rRNa
	79.376	18S, 5.8S & 28S rRNa

Bands 1-4, 6, 9, 13, 14, and 17-20 are shown to be false positives by dot blot analysis and, therefore, are not sequenced. Derived from Rockett et al. (1997). It should be noted that the above genes do not represent the complete spectrum of genes which are up-regulated in rate liver by phenobarbital, but simply represents the genes sequenced and identified to date.

Table 3. Genes down-regulated in rat liver following 3-day exposure to phenobarbital.

Band number (approximate size in bp)	Highest sequence similarity	FASTA-EMBL gene identification
1 (1500) 2 (1200) 3 (1000) 7 (700) 8 (650) 9 (600) 10 (550) 11 (525) 12 (375) 13 (23) 14 (170) 15 (140) Others: (300)	95.3% 92.3% 91.7% Clone 1 77.2% Clone 2 94.5% Clone 3 91.0% Clone 1 86.9% Clone 2 96.2% Clone 2 82.0% 73.8% 95.7% 100.0% Clone 1 97.2% Clone 2 100.0% Clone 3 100.0% Clone 3 100.0% 96.0% 97.3% 96.7%	3-oxoacyl-CoA thiolase Hemopoxin mRNA Alpha-2u-globulin mRNA M.musculus Cl inhibitor Electron transfer flavoprotein M. musculus Topoisomerase 1 (Topo 1), Soares 2NbMT M. musculus (EST) Alpha-2u-globulin (s-type) mRNA Soares mouse NML M. musculus (EST) Soares p3NMF 19.5 M. musculus (EST) Soares mouse NML M. musculus (EST) NCl-CGAP-Pr1 H. sapiens (EST) Ribosomal protein Soares mouse embryo NbME135 (EST) Fibrinogen B-beta-chain Apolipoprotein E gene Soares p3NMF19.5 M. musculus (EST) Stratagene mouse testis (EST) R. norvegicus RASP 1 mRNA
(275)	93.1%	Soares mouse mammary gland (EST)

EST = Expressed sequence tag. Bands 4-6 were shown to be false positives by dot blot analysis and, therefore, were not sequenced. Derived from Rockett et al. (1997). It should be noted that the above genes do not represent the complete spectrum of genes which are down-regulated in rat liver by phenobarbital, but simiply represents the genes sequenced and identified to date.

display' (DD). In this method, all the mRNA species in the control and treated cell populations are amplified in separate reactions using reverse transcriptase-PCR (RT-PCR). The products are then run side-by-side on sequencing gels. Those bands which are present in one display only, or which are much more intense in one

display compared to the other, are differentially expressed and may be recovered for further characterization. One advantage of this system is the speed with which it can be carried out—2 days to obtain a display and as little as a week to make and identify clones.

Two commonly used variations are based on different methods of priming the reverse transcription step (figure 8). One is to use an oligo dT with a 2-base 'anchor' at the 3'-end, e.g. 5'1(dT11)CA 3' (Liang and Pardee 1992). Alternatively, an arbitrary primer may be used for 1st strand cDNA synthesis (Welsh et al. 1992). This variant of RNA fingerprinting has also been called 'RAP' (RNA Arbitrarily Primed)-PCR. One advantage of this second approach is that PCR products may be derived from anywhere in the RNA, including open reading frames. In addition, it can be used for mRNAs that are not polyadenylated, such as many bacterial mRNAs (Wong and McClelland 1994). In both cases, following reverse transcription and denaturation, second strand cDNA synthesis is carried out with an arbitrary primer (arbitrary primers have a single base at each position, as compared to random primers, which contain a mixture of all four bases at each position). The resulting PCR, thus, produces a series of products which, depending on the system (primer length and composition, polymerase and gel system), usually includes 50-100 products per primer set (Band and Sager 1989). When a combination of different dT-anchors and arbitrary primers are used, almost all mRNA species from a cell can be amplified. When the cDNA products from two different populations are analysed side by side on a polyacrylamide gel, differences in expression can be identified and the appropriate bands recovered for cloning and further analysis.

Although DD is perhaps the most popular approach used today for identifying differentially expressed genes, it does suffer from several perceived disadvantages:

- (1) It may have a strong bias towards high copy number mRNAs (Bertioli et al. 1995), although this has been disputed (Wan et al. 1996) and the isolation of very low abundance genes may be achieved in certain circumstances (Guimeraes et al. 1995a).
- (2) The cDNAs obtained often only represent the extreme 3' end of the mRNA (often the 3'-untranslated region), although this may not always be the case (Guimeraes et al. 1995a). Since the 3' end is often not included in Genbank and shows variation between organisms, cDNAs identified by DD cannot always be matched with their genes, even if they have been identified.
- (3) The pattern of differential expression seen on the display often cannot be reproduced on Northern blots, with false positives arising in up to 70% of cases (Sun et al. 1994). Some adaptations have been shown to reduce false positives, including the use of two reverse transcriptases (Sung and Denman 1997), comparison of uninduced and induced cells over a time course (Burn et al. 1994) and comparison of DDPCR-products from two uninduced and two induced lines (Sompayrac et al. 1995). The latter authors also reported that the use of cytoplasmic RNA rather then total RNA reduces false positives arising from nuclear RNA that is not transported to the cytoplasm.

Further details of the background, strengths and weaknesses of the DD technique can be obtained from a review by McClelland et al. (1996) and from articles by Liang et al. (1995) and Wan et al. (1996).

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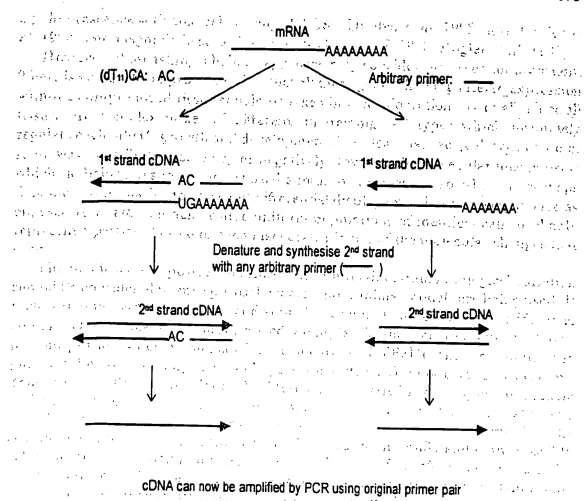


Figure 8. Two approaches to differential display (DD) analysis. 1st strand synthesis can be carried out either with a polydT₁₁ NN primer (where N = G, C or A) or with an arbitrary primer. The use of different combinations of G, C and A to anchor the first strand polydT primer enables the priming of the majority of polyadenylated mRNAs. Arbitrary primers may hybridize at none, one or more places along the length of the mRNA, allowing 1st strand cDNA synthesis to occur at none, one or more points in the same gene. In both cases, 2nd strand synthesis is carried out with an arbitrary primer. Since these arbitrary primers for the 2nd strand may also hybridize to the 1st strand cDNA in a number of different places, several different 2nd strand products may be obtained from one binding point of the 1st strand primer. Following 2nd strand synthesis, the original set of primers is used to amplify the second strand products, with the result that numerous gene sequences are amplified.

Restriction endonuclease-facilitated analysis of gene expression

Serial Analysis of Gene Expression (SAGE)

A more recent development in the field of differential display is SAGE analysis (Velculescu et al. 1995). This method uses a different approach to those discussed so far and is based on two principles. Firstly, in more than 95% of cases, short nucleotide sequences ('tags') of only nine or 10 base pairs provide sufficient information to identify their gene of origin. Secondly, concatonation (linking together in a series) of these tags allows sequencing of multiple cDNAs within a single clone. Figure 9 shows a schematic representation of the SAGE process. In this procedure, double stranded cDNA from the test cells is synthesized with a biotinylated polydT primer. Following digestion with a commonly cutting (4bp recognition sequence) restriction enzyme ('anchoring enzyme'), the 3' ends of the cDNA population are captured with streptavidin beads. The captured population is

split into two and different adaptors ligated to the 5'ends of each group. Incorporated into the adaptors is a recognition sequence for a type IIS restriction enzyme—one which cuts DNA at a defined distance (< 20 bp) from its recognition sequence. Hence, following digestion of each captured cDNA population with the IIS enzyme, the adaptors plus a short piece of the captured cDNA are released. The two populations are then ligated and the products amplified. The amplified products are cleaved with the original anchoring enzyme, religated (concatomers are formed in the process) and cloned. The advantage of this system is that hundreds of gene tags can be identified by sequencing only a few clones. Furthermore, the number of times a given transcript is identified is a quantitative measurement of that gene's abundance in the original population, a feature which facilitates identification of differentially expressed genes in different cell populations.

Some disadvantages of SAGE analysis include the technical difficulty of the method, a large amount of accurate sequencing is required, biased towards abundant mRNAs, has not been validated in the pharmaco/toxicogenomic setting and has only been used to examine well known tissue differences to date.

Gene Expression Fingerprinting (GEF) with a large way to the second of t

A different capture/restriction digest approach for isolating differentially expressed genes has been described by Ivanova and Belyavsky (1995). In this method, RNA is converted to cDNA using biotinylated oligo(dT) primers. The cDNA population is then digested with a specific endonuclease and captured with magnetic streptavidin microbeads to facilitate removal of the unwanted 5' digestion products. The use of restricted 3'-ends alone serves to reduce the complexity of the cDNA fragment pool and helps to ensure that each RNA species is represented by not more than one restriction product. An adaptor is ligated to facilitate subsequent amplification of the captured population. PCR is carried out with one adaptorspecific and one biotinylated polydT primer. The reamplified population is recaptured and the non-biotinylated strands removed by alkaline dissociation. The non-biotinylated strand is then resynthesized using a different adaptor-specific primer in the presence of a radiolabelled dNTP. The labelled immobilized 3'cDNA ends are next sequentially treated with a series of different restriction endonucleases and the products from each digestion analysed by PAGE. The result is a fingerprint composed of a number of ladders (equal to the number of sequential digests used). By comparing test versus control fingerprints, it is possible to identify differentially expressed products which can then be isolated from the gel and cloned. The advantages of this procedure are that it is very robust and reproducible, and the authors estimate that 80-93% of cDNA molecules are involved in the final fingerprint. The disadvantage is that polyacrylamide gels can rarely resolve more than 300-400 bands, which compares poorly to the 1000 or more which are estimated to be produced in an average experiment. The use of 2-D gels such as those described by Uitterlinden et al. (1989) and Hatada et al. (1991) may help to overcome this problem.

A similar method for displaying restriction endonuclease fragments was later described by Prashar and Weissman (1996). However, instead of sequential digestion of the immobolized 3'-terminal cDNA fragments, these authors simply compared the profiles of the control and treated populations without further manipulation.

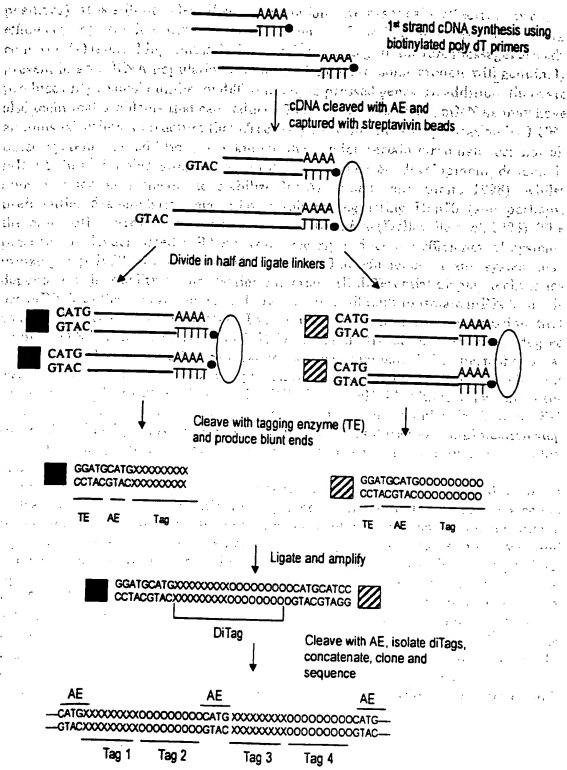


Figure 9. Serial analysis of gene expression (SAGE) analysis. cDNA is cleaved with an anchoring enzyme (AE) and the 3'ends captured using streptavidin beads. The cDNA pool is divided in half and each portion ligated to a different linker, each containing a type IIS restriction site (tagging enzyme, TE). Restriction with the type IIS enzyme releases the linker plus a short length of cDNA (XXXXX and OOOOO indicate nucleotides of different tags). The two pools of tags are then ligated and amplified using linker-specific primers. Following PCR, the products are cleaved with the AE and the ditags isolated from the linkers using PAGE. The ditags are then ligated (during which process, concatenization occurs) and cloned into a vector of choice for sequencing. After Velculescu et al. (1995), with permission.

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DNA arrays (Action 1) to comment of The state of the state of the state of the state of 'Open' differential display systems are cumbersome in that it takes a great deal of time to extract and identify candidate genes and then confirm that they are indeed up- or down-regulated in the treated compared to the control tissue. Normally, the latter process is carried out using Northern blotting or RT-PCR. Even so, each of the aforementioned steps produce a bottleneck to the ultimate goal of rapid analysis of gene expression. These problems will likely be addressed by the development of so-called DNA arrays (e.g. Gress et al. 1992, Zhao et al. 1995, Schena et al. 1996), the introduction of which has signalled the next era in differential gene expression analysis. DNA arrays consist of a gridded membrane or glass chips' containing hundreds or thousands of DNA spots, each consisting of multiple copies of part of a known gene. The genes are often selected based on previously proven involvement in oncogenesis, cell cycling, DNA repair, development and other cellular processes. They are usually chosen to be as specific as possible for each gene and animal species. Human and mouse arrays are already commercially available and a few companies will construct a personalized array to order, for example Clontech Laboratories and Research Genetics Inc. The technique is rapid in that hundreds or even thousands of genes can be spotted on a single array, and that mRNA/cDNA from the test populations can be labelled and used directly as probe. When analysed with appropriate hardware and software, arrays offer a rapid and quantitative means to assess differences in gene expression between two cell populations. Of course, there can only be identification and quantitation of those genes which are in the array (hence the term 'closed' system). Therefore, one approach to elucidating the molecular mechanisms involved in a particular disease/development system may be to combine an open and closed system-a DNA array to directly identify and quantitate the expression of known genes in mRNA populations, and an open system such as SSH to isolate unknown genes which are differentially expressed.

One of the main advantages of DNA arrays is the huge number of gene fragments which can be put on a membrane—some companies have reported gridding up to 60 000 spots on a single glass 'chip' (microscope slide). These high density chipbased micro-arrays will probably become available as mass-produced off-the-shelf items in the near future. This should facilitate the more rapid determination of differential expression in time and dose-response experiments. Aside from their high cost and the technical complexities involved in producing and probing DNA arrays, the main problem which remains, especially with the newer micro-array (gene-chip) technologies, is that results are often not wholly reproducible between arrays. However, this problem is being addressed and should be resolved within the next few years.

EST databases as a means to identify differentially expressed genes.

Expressed sequence tags (ESTs) are partial sequences of clones obtained from cDNA libraries. Even though most ESTs have no formal identity (putative identification is the best to be hoped for), they have proven to be a rapid and efficient means of discovering new genes and can be used to generate profiles of geneexpression in specific cells. Since they were first described by Adams et al. (1991), there has been a huge explosion in EST production and it is estimated that there are now well over a million such sequences in the public domain, representing over half

of all human genes (Hillier et al. 1996). This large number of freely available sequences (both sequence information and clones are normally available royalty-free from the originators) has enabled the development of a new approach towards differential gene expression analysis as described by Vasmatzis et al. (1998). The approach is simple in theory: EST databases are first searched for genes that have a number of related EST sequences from the target tissue of choice, but none or few from non-target tissue libraries. Programmes to assist in the assembly of such sets of overlapping data may be developed in-house or obtained privately or from the internet. For example, the Institute for Genomic Research (TIGR, found at http://www.tigr.org/provides many software tools free of charge to the scientific community. Included amongst these is the TIGR assembler (Sutton et al. 1995), a tool for the assembly of large sets of overlapping data such as ESTs, bacterial artificial chromosomes (BAC)s, or small genomes. Candidate EST clones representing different genes are then analysed using RNA blot methods for size and tissue specificity and, if required, used as probes to isolate and identify the full length cDNA clone for further characterization. In practice however, the method is rather more involved, requiring bioinformatic and computer analysis coupled with confirmatory molecular studies. Vasmatzis et al. (1998) have described several problems in this fledgling approach, such as separating highly homologous sequences derived from different genes and an overemphasis of specificity for some EST sequences. However, since these problems will largely be addressed by the development of more suitable computer algorithms and an increased completeness of the EST database, it is likely that this approach to identifying differentially expressed genes may enjoy more patronage in the future.

Problems and potential of differential expression techniques

The holistic or single cell approach?

When working with in vivo models of differential expression, one of the first issues to consider must be the presence of multiple cell types in any given specimen. For example, a liver sample is likely to contain not only hepatocytes, but also (potentially) Ito cells, bile ductule cells, endothelial cells, various immune cells (e.g. lymphocytes, macrophages and Kupffer cells) and fibroblasts. Other tissues will each have their own distinctive cell populations. Also, in the case of neoplastic tissue, there are almost always normal, hyperplastic and/or dysplastic cells present in a sample. One must, therefore, be aware that genes obtained from a differential display experiment performed on an animal tissue model may not necessarily arise exclusively from the intended 'target' cells, e.g. hepatocytes/neoplastic cells. If appropriate, further analyses using immunohistochemistry, in situ hybridization or in situ RT-PCR should be used to confirm which cell types are expressing the gene(s) of interest. This problem is probably most acute for those studying the differential expression of genes in the development of different cell types, where there is a need to examine homologous cell populations. The problem is now being addressed at the National Cancer Institute (Bethesda, MD, USA) where new microdisection techniques have been employed to assist in their gene analysis programme, the Cancer Genome Anatomy Project (CGAP) (For more information see web site: http://www.ncbi.nlm.nih.gov/ncicgap/intro.html). There are also separation techniques available that utilise cell-specific antigens as a means to isolate target cells,

e.g. fluorescence activated cell sorting (FACS) (Dunbar et al. 1998, Kas-Deelen et al. 1998) and magnetic bead technology (Richard et al. 1998, Rogler et al. 1998).

However, those taking a holistic approach may consider this issue unimportant. There is an equally appropriate view that all those genes showing altered expression within a compromized tissue should be taken into consideration. After all, since all tissues are complex mixes of different, interacting cell types which intimately regulate each other's growth and development, it is clear that each cell type could in some way contribute (positively or negatively) towards the molecular mechanisms which lie behind responses to external stimuli or neoplastic growth. It is perhaps then more informative to carry out differential display experiments using in vivo as opposed to in vitro models, where uniform populations of identical cells probably represent a partial, skewed or even inaccurate picture of the molecular changes that occur.

The incidence and possible implications of inter-individual biological variation should be considered in any approach where whole animal models are being used. It is clear that individuals (humans and animals) respond in different ways to identical stimuli. One of the best characterized examples is the debrisoquine oxidation polymorphism, which is mediated by cytochrome CYP2D6 and determines the pharmacokinetics of many commonly prescribed drugs (Lennard 1993, Meyer and Zanger 1997). The reasons for such differences are varied and complex, but allelic variations, regulatory region polymorphisms and even physical and mental health can all contribute to observed differences in individual responses. Careful thought should, therefore, be given to the specific objectives of the study and to the possible value of pooling starting material (tissue/mRNA). The effect of this can be beneficial through the ironing out of exaggerated responses and unimportant minor fluctuations of (mechanistically) irrelevant genes in individual animals, thus providing a clearer overall picture of the general molecular mechanisms of the response. However, at the same time such minor variations may be of utmost importance in deciding the ability of individual animals to succumb to or resist the effects of a given chemical/disease.

How efficient are differential expression techniques at recovering a high percentage of differentially expressed genes?

A number of groups have produced experimental data suggesting that mammalian cells produce between 8000-15000 different mRNA species at any one time (Mechler and Rabbitts 1981, Hedrick et al. 1984, Bravo 1990), although figures as high as 20-30000 have also been quoted (Axel et al. 1976). Hedrick et al. (1984) provided evidence suggesting that the majority of these belong to the rare abundance class. A breakdown of this abundance distribution is shown in table 1.

When the results of differential display experiments have been compared with data obtained previously using other methods, it is apparent that not all differentially expressed mRNAs are represented in the final display. In particular, rare messages (which, importantly, often include regulatory proteins) are not easily recovered using differential display systems. This is a major shortcoming, as the majority of mRNA species exist at levels of less than 0.005% of the total population (table 1). Bertioli et al. (1995) examined the efficiency of DD templates (heterogeneous mRNA populations) for recovering rare messages and were unable to detect mRNA

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species present at less than 1.2% of the total mRNA population—equivalent to an intermediate or abundant species. Interestingly, when simple model systems (single target only) were used instead of a heterogeneous mRNA population, the same primers could detect levels of target mRNA down to 10000× smaller. These results are probably best explained by competition for substrates from the many PCR products produced in a DD reaction.

The numbers of differentially expressed mRNAs reported in the literature using various model systems provides further evidence that many differentially expressed mRNAs are not recovered. For example, DeRisi et al. (1997) used DNA array technology to examine gene expression in yeast following exhaustion of sugar in the medium, and found that more than 1700 genes showed a change in expression of at least 2-fold. In light of such a finding, it would not be unreasonable to suggest that of the 8000-15 000 different mRNA species produced by any given mammalian cell, up to 1000 or more may show altered expression following chemical stimulation. Whilst this may be an extreme figure, it is known that at least 100 genes are activated /upregulated in Jurkat (T-) cells following IL-2 stimulation (Ullman et al. 1990). In addition, Wan et al. (1996) estimated that interferon-y-stimulated HeLa cells differentially express up to 433 genes (assuming 24000 distinct mRNAs expressed by the cells). However, there have been few publications documenting anywhere near the recovery of these numbers. For example, in using DD to compare normal and regenerating mouse liver, Bauer et al. (1993) found only 70 of 38000 total bands to be different. Of these, 50% (35 genes) were shown to correspond to differentially expressed bands. Chen et al. (1996) reported 10 genes upregulated in female rat liver following ethinyl estradiol treatment. McKenzie and Drake (1997). identified 14 different gene products whose expression was altered by phorbol myristate acetate (PMA, a tumour promoter agent) stimulation of a human myelomonocytic cell line. Kilty and Vickers (1997) identified 10 different gene products whose expression was upregulated in the peripheral blood leukocytes of allergic disease sufferers. Linskens et al. (1995) found 23 genes differentially expressed between young and senescent fibroblasts. Techniques other than DD have also provided an apparent paucity of differentially expressed genes. Using SH for example, Cao et al. (1997) found 15 genes differentially expressed in colorectal cancer compared to normal mucosal epithelium. Fitzpatrick et al. (1995) isolated 17 genes upregulated in rat liver following treatment with the peroxisome proliferator, clofibrate; Philips et al. (1990) isolated 12 cDNA clones which were upregulated in highly metastatic mammary adenocarcinoma cell lines compared to poorly metastatic ones. Prashar and Weissman (1996) used 3' restriction fragment analysis and identified approximately 40 genes showing altered expression within 4 h of activation of Jurkat T-cells. Groenink and Leegwater (1996) analysed 27 gene fragments isolated using SSH of delayed early response phase of liver regeneration and found only 12 to be upregulated.

In the laboratory, SSH was used to isolate up to 70 candidate genes which appear to show altered expression in guinea pig liver following short-term treatment with the peroxisome proliferator, WY-14,643 (Rockett, Swales, Esdaile and Gibson, unpublished observations). However, these findings have still to be confirmed by analysis of the extracted tissue mRNA for differential expression of these sequences.

Whilst the latest differential display technologies are purported to include design and experimental modifications to overcome this lack of efficiency (in both the total number of differentially expressed genes recovered and the percentage that are true

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positives), it is still not clear if such adaptations are practically effective-proving efficiency by spiking with a known amount of limited numbers of artificial rconstruct(s) is one thing, but isolating a high percentage of the rare messages already present in an mRNA population is another. Of course, some models will genuinely produce only a small number of differentially expressed genes. In addition, there are also technical problems that can reduce efficiency. For example, mRNAs may have an unusual primary structure that effectively prevents their amplification by PCRbased systems. In addition, it is known that under certain circumstances not all mRNAs have 3' polyA sites. For example, during Xenopus development, deadenylation is used as a means to stabilize RNAs (Voeltz and Steitz 1998), whilst preferential deadenylation may play a role in regulating Hsp70 (and perhaps, therefore, other stress protein) expression in Drosophila (Dellavalle et al. 1994). The presence of deadenylated mRNAs would clearly reduce the efficiency of systems utilizing a polydT reverse transcription step. The efficiency of any system also depends on the quality of the starting material. All differential display techniques use mRNA as their target material. However, it is difficult to isolate mRNA that is completely free of ribosomal RNA. Even if polydT primers are used to prime first strand cDNA synthesis, ribosomal RNA is often transcribed to some degree (Clontech PCR-Select cDNA Subtraction kit user manual). It has been shown, at least in the case of SSH, that a high rRNA: mRNA ratio can lead to inefficient subtractive hybridization (Clontech PCR-Select cDNA Subtraction kit user manual), and there is no reason to suppose that it will not do likewise in other SH approaches. Finally, those techniques that utilise a presubtraction amplification step (e.g. RDA) may present a skewed representation since some sequences amplify better than others.

Of course, probably the most important consideration is the temporal factor. It is clear that any given differential display experiment can only interrogate a cell at one point in time. It may well be that a high percentage of the genes showing altered expression at that time are obtained. However, given that disease processes and responses to environmental stimuli involve dynamic cascades of signalling, regulation, production and action, it is clear that all those genes which are switched on off at different times will not be recovered and, therefore, vital information may well be missed. It is, therefore, imperative to obtain as much information about the model system beforehand as possible, from which a strategy can be derived for targeting specific time points or events that are of particular interest to the investigator. One way of getting round this problem of single time point analysis is to conduct the experiment over a suitable time course which, of course, adds substantially to the amount of work involved.

How sensitive are differential expression technologies?

There has been little published data that addresses the issue of how large the change in expression must be for it to permit isolation of the gene in question with the various differential expression technologies. Although the isolation of genes whose expression is changed as little as 1.5-fold has been reported using SSH (Groenink and Leegwater 1996), it appears that those demonstrating a change in excess of 5-fold are more likely to be picked up. Thus, there is a 'grey zone' in between where small changes could fade in and out of isolation between

experiments and animals. DD, on the other hand, is not subject to this grey zone since, unlike SH approaches, it does not amplify the difference in expression between two sampless Wan et al. (1996) reported that differences in expression of twofold or more are detectable using DD.

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Resolution and visualization of differential expression products

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It seems highly improbable with current technology that a gel system could be developed that is able to resolve all gene species showing altered expression in any given test system (be it SH-or DD-based). Polyacrylamide gel electrophoresis (PAGE) can resolve size differences down to 0.2% (Sambrook et al. 1989) and are used as standard in DD experiments. Even so, it is clear that a complex series of gene products such as those seen in a DD will contain unresolvable components. Thus, what appears to be one band in a gel may in fact turn out to be several. Indeed, it has been well documented (Mathieu-Daude et al. 1996, Smith et al. 1997) that a single band extracted from a DD often represents a composite of heterogeneous products, and the same has been found for SSH displays in this laboratory (Rockett et al. 1997). One possible solution was offered by Mathieu-Daude et al. (1996), who extracted and reamplified candidate bands from a DD display and used single strand conformation polymorphism (SSCP) analysis to confirm which components represented the truly differentially expressed product.

Many scientists often try to avoid the use of PAGE where possible because it is technically more demanding than agarose gel electrophoresis (AGE). Unfortunately, high resolution agarose gels such as Metaphor (FMC, Lichfield, UK) and AquaPor HR (National Diagnostics, Hessle, UK), whilst easier to prepare and manipulate than PAGE, can only separate DNA sequences which differ in size by around 1.5-2% (15-20 base pairs for a 1Kb fragment). Thus, SSH, RDA or other such products which differ in size by less than this amount are normally not resolvable. However, a simple technique does in fact exist for increasing the resolving power of AGE—the inclusion of HA-red (10-phenyl neutral red-PEG ligand) or HA-yellow (bisbenzamide-PEG ligand) (Hanse Analytik GmbH, Bremen, Germany) in a gel separates identical or closely sized products on base content. Specifically, HA-red and -yellow selectively bind to GC and AT DNA motifs, respectively (Wawer et al. 1995, Hanse Analytik 1997, personal communication). Since both HA-stains possess an overall positive charge, they migrate towards the cathode when an electric field is applied. This is in direct opposition to DNA, which is negatively charged and, therefore, migrates towards the anode. Thus, if two DNA clones are identical in size (as perceived on a standard high resolution agarose gel), but differ in AT/GC content, inclusion of a HA-dye in the gel will effectively retard the migration of one of the sequences compared to the other, effectively making it apparently larger and, thus, providing a means of differentiating between the two. The use of HA-red has been shown to resolve sequences with an AT variation of less than 1% (Wawer et al. 1995), whilst Hanse Analytik have reported that HA staining is so sensitive that in one case it was used to distinguish two 567bp sequences which differed by only a single point mutation (Hanse Analytik 1996, personal communication). Therefore, if one wishes to check whether all the clones produced from a specific band in a differential display experiment are derived from the same gene species, a small amount of reamplified or digested clone can be run on a standard high resolution gel, and a second aliquot

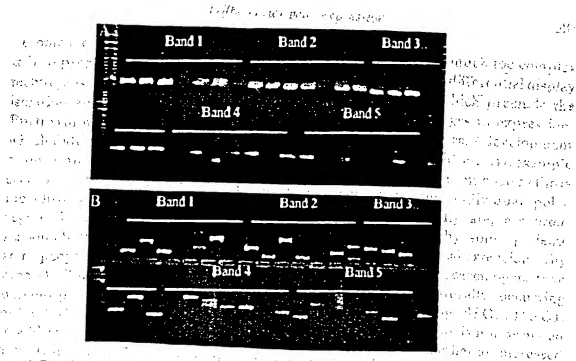


Figure 10. Discrimination of clones of identical/nearly identical size using HA-red. Bands of decreasing size (1-5) were extracted from the final display of a suppression subtractive hybridization experiment and cloned. Seven colonies were picked at random from each cloned band and their inserts amplified using PCR. The products were run on two gels, (A) a high resolution 2% agarose gel, and (B) a high resolution 2% agarose gel containing 1 U/ml HA-red. With few exceptions, all the clones from each band appear to be the same size (gel A). However, the presence of HA-red (gel B), which separates identically-sized DNA fragments based on the percentage of GC within the sequence, clearly indicates the presence of different gene species within each band. For example, even though all five re-amplified clones of band 1 appear to be the same size, at least four different gene species are represented.

in a similar gel containing one of the HA-stains. The standard gel should indicate any gross size differences, whilst the HA-stained gel should separate otherwise unresolvable species (on standard AGE) according to their base content. Geisinger et al. (1997) reported successful use of this approach for identifying DD-derived clones. Figure 10 shows such an experiment carried out in this laboratory on clones obtained from a band extracted from an SSH display.

An alternative approach is to carry out a 2-D analysis of the differential display products. In this approach, size-based separation is first carried out in a standard agarose gel. The gel slice containing the display is then extracted and incorporated in to a HA gel for resolution based on AT/GC content.

Of course, one should always consider the possibility of there being different gene species which are the same size and have the same GC/AT content. However, even these species are not unresolvable given some effort—again, one might use SSCP, or perhaps a denaturing gradient gel electrophoresis (DGGE) or temperature gradient field electrophoresis (TGGE) approach to resolve the contents of a band, either directly on the extracted band (Suzuki et al. 1991) or on the reamplified product.

The requirement of some differential display techniques to visualize large numbers of products (e.g. DD and GEF) can also present a problem in that, in terms of numbers, the resolution of PAGE rarely exceeds 300-400 bands. One approach to overcoming this might be to use 2-D gels such as those described by Uitterlinden et al. (1989) and Hatada et al. (1991).

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Extraction of differentially expressed bands from a gel can be complex since, in some cases (e.g. DD, GEF), the results are visualized by autoradiographic means, such that precise overlay of the developed film on the gel must occur if the correct band is to be extracted for further analysis. Clearly, a misjudged extraction can account for many man-hours lost. This problem, and that of the use of radioisotopes, has been addressed by several groups. For example, Lohmann et al. (1995) demonstrated that silver staining can be used directly to visualize DD bands in horizontal PAGs. An et al. (1996) avoided the use of radioisotopes by transferring a small amount (20-30%) of the DNA from their DD to a nylon membrane, and visualizing the bands using chemiluminescent staining before going back to extract the remaining DNA from the gel. Chen and Peck (1996) went one step further and transferred the entire DD to a nylon membrane. The DNA bands were then visualized using a digoxigenin (DIG) system (DIG was attached to the polydT primers used in the differential display procedure). Differentially expressed bands were cut from the membrane and the DNA eluted by washing with PCR buffer prior to reamplification.

One of the advantages of using techniques such as SSH and RDA is that the final display can be run on an agarose gel and the bands visualized with simple ethidium bromide staining. Whilst this approach can provide acceptable results, overstaining with SYBR Green I or SYBR Gold nucleic acid stains (FMC) effectively enhances the intensity and sharpness of the bands. This greatly aids in their precise extraction and often reveals some faint products that may otherwise be overlooked. Whilst differential displays stained with SYBR Green I are better visualized using short wavelength UV (254 nm) rather than medium wavelength (306 nm), the shorter wavelength is much more DNA damaging. In practice, it takes only a few seconds to damage DNA extracted under 254 nm irradiation, effectively preventing reamplification and cloning. The best approach is to overstain with SYBR Green I and extract bands under a medium wavelength UV transillumination.

The possible use of 'microfingerprinting' to reduce complexity

Given the sheer number of gene products and the possible complexity of each band, an alternative approach to rapid characterization may be to use an enhanced analysis of a small section of a differential display—a 'sub-fingerprint' or 'microfingerprint'. In this case, one could concentrate on those bands which only appear in a particular chosen size region. Reducing the fingerprint in this way has at least two advantages. One is that it should be possible to use different gel types, concentrations and run times tailored exactly to that region. Currently, one might run products from 100-3000 + bp on the same gel, which leads to compromize in the gel system being used and consequently to suboptimal resolution, both in terms of size and numbers, and can lead to problems in the accurate excision of individual bands. Secondly, it may be possible to enhance resolution by using a 2-D analysis using a HA-stain, as described earlier. In summary, if a range of gene product sizes is carefully chosen to included certain 'relevant' genes, the 2-D system standardized, and appropriate gene analysis used, it may be possible to develop a method for the early and rapid identification of compounds which have similar or widely different cellular effects. If the prognosis for exposure to one or more other chemicals which display a similar profile is already known, then one could perhaps predict similar effects for any new compounds which show a similar micro-fingerprint.

An alternative approach to microfingerprinting is to examine altered expression in specific families of genes through careful selection of PCR primers and for postreaction analysis. Stress genes, growth factors and/or their receptors, cell cycling genes, cytochromes P450 and regulatory proteins might be considered as candidates for analysis in this way. Indeed, some off-the-shelf DNA arrays (e.g. Clontech's Atlas cDNA Expression Array series) already anticipated this to some degree by grouping together genes involved in different responses e.g. apoptosis, stress, DNAdamage response etc. Carlot Branch will be the Conference where $\mathcal{F}(x)$ is the specific property of (x,y)

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False positives

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The product of the state of the The generation of false positives has been discussed at length amongst the differential display community (Liang et al. 1993, 1995, Nishio et al. 1994, Sun et al. 1994, Sompayrac et al. 1995). The reason for false positives varies with the technique being used. For instance, in RDA, the use of adaptors which have not been HPLC purified can lead to the production of false positives through illegitimate ligation events (O'Neill and Sinclair 1997), whilst in DD they can arise through PCR artifacts and illegitemate transcription of rRNA. In SH, false positives appear to be derived largely from abundant gene species, although some may arise from cDNA/mRNA species which do not undergo hybridization for technical reasons.

A quick screening of putative differentially expressed clones can be carried out using a simple dot blot approach, in which labelled first strand probes synthesized from tester and driver mRNA are hybridized to an array of said clones (Hedrick et al. 1984, Sakaguchi et al. 1986). Differentially expressed clones will hybridize to tester probe, but not driver. The disadvantage of this approach is that rare species may not generate detectable hybridization signals. One option for those using SSH is to screen the clones using a labelled probe generated from the subtracted cDNA from which it was derived, and with a probe made from the reverse subtraction reaction (ClonTechniques 1997a). Since the SSH method enriches rare sequences, it should be possible to confirm the presence of clones representing low abundance genes. Despite this quick screening step, there is still the need to go back to the original mRNA and confirm the altered expression using a more quantitative approach. Although this may be achieved using Northern blots, the sensitivity is poor by today's high standards and one must rely on PCR methods for accurate and sensitive determinations (see below).

Sequence analysis

The majority of differential display procedures produce final products which are between 100 and 1000bp in size. However, this may considerably reduce the size of the sequence for analysis of the DNA databases. This in turn leads to a reduced confidence in the result-several families of genes have members whose DNA sequences are almost identical except in a few key stretches, e.g. the cytochrome P450 gene superfamily (Nelson et al. 1996). Thus, does the clone identified as being almost identical to gene X₀ really come from that gene, or its brother gene X₁ or its as yet undiscovered sister X2? For example, using SSH, part of a gene was isolated,

which was up-regulated in the liver of rats exposed to Wy-14,643 and was identified by a FASTA search as being transferrin (data not shown). However, transferrin is known to be downregulated by hypolipidemic peroxisome proliferators such as Wy-14,643 (Hertz et al. 1996), and this was confirmed with subsequent RT-PCR analysis. This suggests that the gene sequence isolated may belong to a gene which is closely related to transferrin, but is regulated by a different mechanism.

A further problem associated with SH technology is redundancy. In most cases before SH is carried out, the cDNA population must first be simplified by restriction

digestion. This is important for at least two reasons:

(1) To reduce complexity—long cDNA fragments may form complex networks which prevent the formation of appropriate hybrids, especially at the high concentrations required for efficient hybridization.

(2) Cutting the cDNAs into small fragments provides better representation of individual genes. This is because genes derived from related but distinct members of gene families often have similar coding sequences that may crosshybridize and be eliminated during the subtraction procedure (Ko 1990). Furthermore, different fragments from the same cDNA may differ considerably in terms of hybridization and amplification and, thus, may not efficiently do one or the other (Wang and Brown 1991). Thus, some fragments from differentially expressed cDNAs may be eliminated during subtractive hybridization procedures. However, other fragments may be enriched and isolated. As a consequence of this, some genes will be cut one or more times, giving rise to two or more fragments of different sizes. If those same genes are differentially expressed, then two or more of the different size fragments may come through as separate bands on the final differential display, increasing the observed redundancy and increasing the number of redundant sequencing reactions.

Sequence comparisons also throw up another important point—at what degree of sequence similarity does one accept a result. Is 90% identitiy between a gene derived from your model species and another acceptably close? Is 95% between your sequence and one from the same species also acceptable? This problem is particularly relevant when the forward and reverse sequence comparisons give similar sequences with completely different gene species! An arbitrary decision seems to be to allocate genes that are definite (95% and above similarity) and then group those between 60 and 95% as being related or possible homologues.

Quantitative analysis

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At some point, one must give consideration to the quantitative analysis of the candidate genes, either as a means of confirming that they are truly differentially expressed, or in order to establish just what the differences are. Northern blot analysis is a popular approach as it is relatively easy and quick to perform. However, the major drawback with Northern blots is that they are often not sensitive enough to detect rare sequences. Since the majority of messages expressed in a cell are of low abundance (see table 1), this is a major problem. Consequently, RT-PCR may be the method of choice for confirming differential expression. Although the procedure is somewhat more complex than Northern analysis, requiring synthesis of primers and optimization of reaction conditions for each gene species, it is now possible to set up high throughput PCR systems using mulitchannel pipettes, 96 +-well plates and

appropriate thermal cycling technology. Whilst quantitative analysis is more desirable, being more accurate and without reliance on an internal standard, the money and time needed to develop a competitor molecule is often excessive, especially when one might be examining tens or even hundreds of gene species. The use of semi-quantitative analysis is simpler, although still relatively involved. One must first of all choose an internal standard that does not change in the test cells compared to the controls. Numerous reference genes have been tried in the past, for example interferon-gamma (IFN-7, Frye et al. 1989), \(\beta\)-actin (Heuval et al. 1994), glyceraldehyde-3-phosphate dehydrogenase (GAPDH, Wong et al. 1994), dihydrofolate reductase (DHFR, Mohler and Butler 1991), \(\beta\)-2-microglobulin (\(\beta\)-2m, Murphy et al. 1990), hypoxanthine phosphoribosyl transferase (HPRT, Foss et al. 1998) and a number of others (ClonTechniques 1997b). Ideally, an internal standard should not change its level of expression in the cell regardless of cell age, stage in the cell cycle or through the effects of external stimuli. However, it has been shown on numerous occasions that the levels of most housekeeping genes currently used by the research community do in fact change under certain conditions and in different tissues (ClonTechniques 1997b). It is imperative, therefore, that preliminary experiments be carried out on a panel of housekeeping genes to establish their suitability for use in the model system.

Interpretation of quantitative data must also be treated with caution. By comparing the lists of genes identified by differential expression one can perhaps gain insight into why two different species react in different ways to external stimuli. For example, rats and mice appear sensitive to the non-genotoxic effects of a wide range of peroxisome proliferators whilst Syrian hamsters and guinea pigs are largely resistant (Orton et al. 1984, Rodricks and Turnbull 1987, Lake et al. 1989, 1993, Makowska et al. 1992). A simplified approach to resolving the reason(s) why is to compare lists of up- and down-regulated genes in order to identify those which are expressed in only one species and, through background knowledge of the effects of the said gene, might suggest a mechanism of facilitated non-genotoxic carcinogenesis or protection. Of course, the situation is likely to be far more complex. Perhaps if there were one key gene protecting guinea pig from non-genotoxic effects and it was upregulated 50 times by PPs, the same gene might only be up-regulated five times in the rat. However, since both were noted to be upregulated, the importance of the gene may be overlooked. Just to complicate matters, a large change in expression does not necessarily mean a biologically important change. For example, what is the true relevance of gene Y which shows a 50-fold increase after a particular treatment, and gene Z which shows only a 5-fold increase? If one examines the literature one may find that historically, gene Y has often been shown to be up-regulated 40-60fold by a number of unrelated stimuli—in light of this the 50-fold increase would appear less significant. However, the literature may show that gene Z has never been recorded as having more than doubled in expression—which makes your 5-fold increase all the more exciting. Perhaps even more interesting is if that same 5-fold increase has only been seen in related neoplasms or following treatment with related

Problems in using the differential display approach

Differential display technology originally held promise of an easily obtainable 'fingerprint' of those genes which are up- or down-regulated in test animals/cells in a developmental process or following exposure to given stimuli. However, it has

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become clear that the fingerprinting process, whilst still valid, is much too complex to be represented by a single technique profile. This is because all differential display techniques have common and or unique technical problems which preclude the isolation and identification of all those genes which show changes in expression. Furthermore, there are important genetic changes related to disease development which differential expression analysis is simply not designed to address. An example of this is the presence of small deletions, insertions, or point mutations such as those seen in activated oncogenes, tumour suppressor genes and individual polymorphisms. Polymorphic variations, small though they usually are, are often regarded as being of paramount importance in explaining why some patients respond better than others to certain drug treatments (and, in logical extension, why some people are less affected by potentially dangerous xenobiotics/carcinogens than others). The identification of such point mutations and naturally occurring polymorphisms requires the subsequent application of sequencing, SSCP, DGGE or TGGE to the gene of interest. Furthermore, differential display is not designed to address issues such as alternatively spliced gene species of whether an increased abundance of mRNA is a result of increased transcription or increased mRNA

Conclusions

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Perhaps the main advantage of open system differential display techniques is that they are not limited by extant theories or researcher bias in revealing genes which are differentially expressed, since they are designed to amplify all genes which demonstrate altered expression. This means that they are useful for the isolation of previously unknown genes which may turn out be useful biomarkers of a particular state or condition. At least one open system (SAGE) is also quantitative, thus eliminating the need to return to the original mRNA and carry out Northern/PCR analysis to confirm the result. However, the rapid progress of genome mapping projects means that over the next 5-10 years or so, the balance of experimental use will switch from open to closed differential display systems, particularly DNA arrays. Arrays are easier and faster to prepare and use, provide quantitative data, are suitable for high throughput analysis and can be tailored to look at specific signalling pathways or families of genes. Identification of all the gene sequences in human and common laboratory animals combined with improved DNA array technology, means that it will soon no longer be necessary to try to isolate differentially expressed genes using the technically more demanding open system approach. Thus, their main advantage (that of identifying unknown genes) will be largely eradicated. It is likely, therefore, that their sphere of application will be reduced to analysis of the less common laboratory species, since it will be some time yet before the genomes of such animals as zebrafish, electric eels, gerbils, crayfish and squid, for example, will

Of course, in the end the question will always remain: What is the functional/ biological significance of the identified, differentially expressed genes? One persistent problem is understanding whether differentially expressed genes are a cause or consequence of the altered state. Furthermore, many chemicals, such as non-genotoxic carcinogens, are also mitogens and so genes associated with replication will also be upregulated but may have little or nothing to do with the

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that the protection of the solution of the section carcinogenic effect. Whilst differential display technology cannot hope to answer these questions, it does provide a springboard from which identification, regulatory and functional studies can be launched. Understanding the molecular mechanism of cellular responses is almost impossible without knowing the regulation and function of those genes and their condition (e.g. mutated). In an abstract sense, differential display can be likened to a still photograph, showing details of a fixed moment in time. Consider the Historian who knows the outcome of a battle and the placement and condition of the troops before the battle commenced, but is asked to try and deduce how the battle progressed and why it ended as it did from a few still photographs—an impossible task. In order to understand the battle, the Historian must find out the capabilities and motivation of the soldiers and their commanding officers, what the orders were and whether they were obeyed. He must examine the terrain, the remains of the battle and consider the effects the prevailing weather conditions exerted. Likewise, if mechanistic answers are to be forthcoming, the scientist must use differential display in combination with other techniques, such as knockout technology, the analysis of cell signalling pathways, mutation analysis and time and dose response analyses. Although this review has emphasized the importance of differential gene profiling, it should not be considered in isolation and the full impact of this approach will be strengthened if used in combination with functional genomics and proteomics (2-dimensional protein gels from isoelectric focusing and subsequent SDS electrophoresis and virtual 2D-maps using capillary electrophoresis). Proteomics is attracting much recent attention as many of the changes resulting in differential gene expression do not involve changes in mRNA levels, as decribed extensively herein, but rather protein-protein, protein-DNA and protein phosphorylation events which would require functional genomics or proteomic technologies for investigation.

Despite the limitations of differential display technology, it is clear that many potential applications and benefits can be obtained from characterizing the genetic changes that occur in a cell during normal and disease development and in response to chemical or biological insult. In light of functional data, such profiling will provide a 'fingerprint' of each stage of development or response, and in the long term should help in the elucidation of specific and sensitive biomarkers for different types of chemical/biological exposure and disease states. The potential medical and therapeutic benefits of understanding such molecular changes are almost immeasurable. Amongst other things, such fingerprints could indicate the family or even specific type of chemical an individual has been exposed to plus the length and/or acuteness of that exposure, thus indicating the most prudent treatment. They may also help uncover differences in histologically identical cancers, provide diagnostic tests for the earliest stages of neoplasia and, again, perhaps indicate the most efficacious treatment.

The Human Genome Project will be completed early in the next century and the DNA sequence of all the human genes will be known. The continuing development and evolution of differential gene expression technology will ensure that this knowledge contributes fully to the understanding of human disease processes.

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References

- ADAMS, M. D., KELLEY, J. M., GOCAYNE, J. D., DUBNICK, M., POLYMEROPOULOS, M. H., XIAO, H., MERRIL, C. R., WU, A., OLDE, B., MORENO, R. F., KERLAVAGE, A. R., McCombie, W. R. and Ventor, J. C., 1991, Complementary DNA sequencing: expressed sequence tags and human genome project. Science, 252, 1651-1656.
- An, G., Luo, G., Veltri, R. W. and O'Hara, S. M., 1996, Sensitive non-radioactive differential display method using chemiluminescent detection. *Biotechniques*, 20, 342-346.
- AXEL, R., FEIGELSON, P. and SCHULTZ, G., 1976, Analysis of the complexity and diversity of mRNA from chicken liver and oviduct. Cell, 7, 247-254.
- BAND, V. and SAGER, R., 1989, Distinctive traits of normal and tumor-derived human mammary epithelial cells expressed in a medium that supports long-term growth of both cell types. Proceedings of the Naional Academy of Sciences, USA, 86,1249-1253.
- BAUER, D., MULLER, H., REICH, J., RIEDEL, H., AHRENKIEL, V., WARTHOE, P. and STRAUSS, M., 1993, Identification of differentially expressed mRNA species by an improved display technique (DDRT-PCR). Nucleic Acids Research, 21, 4272-4280.
- BERTIOLI, D. J., SCHLICHTER, U. H. A., ADAMS, M. J., BURROWS, P. R., STEINBISS, H.-H. and ANTONIW, J. F., 1995, An analysis of differential display shows a strong bias towards high copy number mRNAs. Nucleic Acids Research, 23, 4520-4523.
- Bravo, R., 1990, Genes induced during the G0/G1 transition in mouse fibroblasts. Seminars in Cancer Biology, 1, 37-46.
- Burn, T. C., Petrovick, M. S., Hohaus, S., Rollins, B. J. and Tenen, D. G., 1994, Monocyte chemoattractant protein-1 gene is expressed in activated neutrophils and retinoic acid-induced human myeloid cell lines. *Blood*, 84, 2776-2783.
- CAO, J., CAI, X., ZHENG, L., GENG, L., SHI, Z., PAO, C. C. and ZHENG, S., 1997, Characterisation of colorectal cancer-related cDNA clones obtained by subtractive hybridisation screening. Journal of Cancer Research and Clinical Oncology, 123, 447-451.
- Cassidy, S. B., 1995, Uniparental disomy and genomic imprinting as causes of human genetic disease. Environmental and Molecular Mutagenesis, 25 (Suppl 26), 13-20.
- CHANG, G. W. and Terzaghi-Howe, M., 1998, Multiple changes in gene expression are associated with normal cell-induced modulation of the neoplastic phenotype. Cancer Research, 58, 4445-4452.
- CHEN, J., SCHWARTZ, D. A., YOUNG, T. A., NORRIS, J. S. and YAGER, J. D., 1996, Identification of genes whose expression is altered during mitosuppression in livers of ethinyl estradiol-treated female rats. Carcinogenesis, 17, 2783-2786.
- CHEN, J. J. W. and PECK, K., 1996, Non-radioactive differential display method to directly visualise and amplify differential bands on nylon membrane. Nucleic Acid Research, 24, 793-794.
- CLON TECHNIQUES, 1997a, PCR-Select Differential Screening Kit—the nextstep after Clontech PCR-Select cDNA subtraction. ClonTechniques, XII, 18-19.
- CLON TECHNIQUES, 1997b, Housekeeping RT-PCR amplimers and cDNA probes. ClonTechniques, XII, 15-16.
- DAVIS, M. M., COHEN, D. I., NIELSEN, E. A., STEINMETZ, M., PAUL, W. E. and Hood, L., 1984, Cell-type-specific cDNA probes and the murine I region: the localization and orientation of Ad alpha. Proceedings of the National Academy of Sciences (USA), 81, 2194-2198.
- Dellavalle, R. P., Peterson, R. and Lindquist, S., 1994, Preferential deadenylation of HSP70 mRNA plays a key role in regulating Hsp70 expression in Drosophila melanogaster. *Molecular and Cell Biology*, 14, 3646-3659.
- DERISI, J. L., VASHWANATH, R. L. and BROWN, P., 1997, Exploring the metabolic and genetic control of gene expression on a genomic scale. Science, 278, 680-686.
- DIATCHENKO, L., LAU, Y.-F. C., CAMPBELL, A. P., CHENCHIK, A., MOOADAM, F., HUANG, B., LUKYANOV, K., GURSKAYA, N., SVERDLOV, E. D. and SIEBERT, P. D., 1996, Suppression subtractive hybridisation: A method for generating differentially regulated or tissue-specific cDNA probes and libraries. Proceedings of the National Academy of Sciences (USA), 93, 6025-6030.
- Dogra, S. C., Whitelaw, M. L. and May, B. K., 1998, Transcriptional activation of cytochrome P450 genes by different classes of chemical inducers. Clinical and Experimental Pharmacology and Physiology, 25, 1-9.
- Duguid, J. R. and Dinauer, M. C., 1990, Library subtraction of in vitro cDNA libraries to identify differentially expressed genes in scrapic infection. Nucleic Acids Research, 18, 2789-2792.
- Dunbar, P. R., Ogg, G. S., Chen, J., Rust, N., van der Bruggen, P. and Cerundolo, V., 1998, Direct isolation, phenotyping and cloning of low-frequency antigen-specific cytotoxic T lymphocytes from peripheral blood. Current Biology, 26, 413-416.

- FITZPATRICK, D. R., GERMAIN LEE, E. and VALLE, D., 1995, Isolation and characterisation of rat and human cDNAs encoding a novel putative peroxisomal enoyl-CoA hydratase. Genomics, 27,
 - Foss, D. L., BAARSCH, M. J. and Murtaugh, M. P., 1998, Regulation of hypoxanthine phosphoribosyltransferase, glyceraldehyde-3-phosphate dehydrogenase and beta-actin mRNA expression in porcine immune cells and tissues. Animal Biotechnology, 9, 67-78.
- FRYE, R. A., BENZ, C. C. and Liu, E., 1989, Detection of amplified oncogenes by differential polymerase chain reaction. Oncogene, 4, 1153-1157.
- GEISINGER, A., RODRIGUEZ, R., ROMERO, V. and WETISTEIN R., 1997, A simple method for screening cDNAs arising from the cloning of RNA differential display bands. Elsevier Trends Journals Technical Tips Online, http://tto.trends.com, document T01110.
- GRESS, T. M., HOHEISEL, J. D., LENNON, G. G., ZEHETNER, G. and LEHRACH, H., 1992, Hybridisation fingerprinting of high density cDNA filter arrays with cDNA pools derived from whole tissues.

 Mammalian Genome, 3, 609-619
- GRIFFIN, G. and KRISHNA, S., 1998, Cytokines in infectious diseases. Journal of the Royal College of
- GROENINK, M. and LEEGWATER, A. C. J., 1996, Isolation of delayed early genes associated with liver regeneration using Clontech PCR-select subtraction technique. Clontechniques, XI, 23-24.
- GUIMARAES, M. J., BAZAN, J. F., ZLOTNIK, A., WILES, M. V., GRIMALDI, J.C., LEE, F. and McClanahan, T., 1995b, A new approach to the study of haematopoietic development in the yolk sac and embryoid bodies. Development, 121, 3335-3346.
- GUIMERAES, M. J., LEE, F., ZLOTNIK, A. and McClanahan, T., 1995a, Differential display by PCR: novel findings and applications. Nucleic Acids Research, 23, 1832-1833.
- GURSKAYA, N. G., DIATCHENKO, L., CHENCHIK, P. D., SIEBERT P.P. D., KHASPEKOV, G. L., LUKYANOV, K. A., VAGNER, L. L., ERMOLAEVA, O. D., LUKYANOV, S. A. and SVERDLOV, E. D., 1996, Equalising cDNA subtraction based on selective suppression of polymerase chain reaction: Cloning of Jurkat cell transcripts induced by phytohemaglutinin and phorbol 12-Myrystate 13-Acetate. Analytical Biochemistry, 240, 90-97.
- HAMPSON, I. N. and HAMPSON, L., 1997, CCLS and DROP—subtractive cloning made easy. Life Science News (A publication of Amersham Life Science), 23, 22-24.
- HAMPSON, I. N., HAMPSON, L. and DEXTER, T. M., 1996, Directional random oligonucleotide primed (DROP) global amplification of cDNA: its application to subtractive cDNA cloning. Nucleic
- HAMPSON, I. N., POPE, L., COWLING, G. J. and DEXTER, T. M., 1992, Chemical cross linking subtraction (CCLS): a new method for the generation of subtractive hybridisation probes. Nucleic Acids Research, 20, 2899
- HARA, E., KATO, T., NAKADA, S., SEKIYA, S. and ODA, K., 1991, Subtractive cDNA cloning using oligo(dT)30-latex and PCR: isolation of cDNA clones specific to undifferentiated human embryonal carcinoma cells. Nucleic Acids Research, 19, 7097-7104.
- HATADA, I., HAYASHIZAKE, Y., HIROTSUNE, S., KOMATSUBARA, H. and MUKAI, T., 1991, A genomic scanning method for higher organisms using restriction sites as landmarks. Proceedings of the National Academy of Sciences (USA), 88, 9523-9527.
- HECHT, N., 1998, Molecular mechanisms of male sperm cell differentiation. Bioessays, 20, 555-561.
- HEDRICK, S., COHEN, D. I., NIELSEN, E. A. and DAVIS, M. E., 1984, Isolation of T cell-specific membrane-associated proteins. Nature, 308, 149-153.
- HERTZ, R., SECKBACH, M., ZAKIN, M. M. and BAR-TANA, J., 1996, Transcriptional suppression of the transferrin gene by hypolipidemic peroxisome proliferators. Journal of Biological Chemistry, 271,
- HEUVAL, J. P. V., CLARK, G. C., KOHN, M. C., TRITSCHER, A. M., GREENLEE, W. F., LUCIER, G. W. and BELL, D. A., 1994, Dioxin-responsive genes: Examination of dose-response relationships using quantitative reverse transciptase-polymerase chain reaction. Cancer Research, 54, 62-68.
- HILLER, L. D., LENNON, G., BECKER, M., BONALDO, M. F., CHIAPELLI, B., CHISSOE, S., DIETRICH, N., DUBUQUE, T., FAVELLO, A., GISH, W., HAWKINS, M., HULTMAN, M., KUCABA, T., LACY, M., LE, M., LE, N., MARDIS, E., MOORE, B., MORRIS, M., PARSONS, J., PRANGE, C., RIFKIN, L., ROHLFING, T., SCHELLENBERG, K., SOARES, M. B., TAN, F., THIERRY-MEG, J., TREVASKIS, E., UNDERWOOD, K., Wohldman, P., Waterston, R., Wilson, R and Marra, M., 1996, Generation and analysis of 280,000 human expressed sequence tags. Genome Research, 6, 807-828.
- HUBANK, M. and SCHATZ, D. G., 1994, Identifying differences in mRNA expression by representational difference analysis. Nucleic Acids Research, 22, 5640-5648.
- HUNTER, T., 1991, Cooperation between oncogenes. Cell, 64, 249-270.
- Ivanova, N. B. and Belyavsky, A. V., 1995, Identification of differentially expressed genes by restriction endonuclease-based gene expression fingerprinting. Nucleic Acids Research, 23, 2954-2958.
- JAMES, B. D. and HIGGINS, S. J., 1985, Nucleic Acid Hybridisation (Oxford: IRL Press Ltd).
- KAS-DEELEN, A. M., HARMSEN, M. C., DE MAAR, E. F. and VAN SON, W. J., 1998, A sensitive method for

- quantifying cytomegalic endothelial cells in peripheral blood from cytomegalovirus-infected patients. Clinical Diagnostic and Laboratory Immunology, 5, 622-626.
- Kilty, I. and Vickers, P., 1997, Fractionating DNA fragments generated by differential display PCR. Strategies Newsletter (Strategies), 10, 50-51.
- KLEINJAN, D.-J. and VAN HEYNINGEN, V., 1998, Position effect in human genetic disease. Human and Molecular Genetics, 7, 1611-1618.
- Ko, M. S.; 1990, An equalized cDNA library by the reassociation of short double-stranded cDNAs.

 Nucleic Acids Research, 18, 5705-5711.
- LAKE, B. G., EVANS, J. G., CUNNINGHAME, M. E. and PRICE, R. J., 1993, Comparison of the hepatic effects of Wy-14,643 on peroxisome proliferation and cell replication in the rat and Syrian hamster. Environmental Health Perspectives, 101, 241-248.
- hamster. Environmental Health Perspectives, 101, 241-248.

 LAKE, B.G., EVANS, J. G., GRAY, T. J. B., KOROSI, S. A. and NORTH, C. J., 1989, Comparative studies of nafenopin-induced hepatic peroxisome proliferation in the rat, Syrian hamster, guiea pig and marmoset. Toxicology and Applied Pharmacology, 99, 148-160.
- LENNARD, M. S., 1993, Genetically determined adverse drug reactions involving metabolism. Drug Safety, 9, 60-77.
- LEWY, S., TODD, S. C. and MAECKER, H. T., 1998, CD81(TAPA-1): a molecule involved in signal transduction and cell adhesion in the immune system. Annual Review of Immunology, 16, 89-109.
- LIANG, P. and PARDEE, A. B., 1992, Differential display of eukaryotic messenger RNA by means of the polymerase chain reaction. Science, 257, 967-971.
- LIANG, P., AVERBOUKH, L., KEYOMARSI, K., SAGER, R. and PARDEE, A., 1992, Differential display and cloning of messenger RNAs from human breast cancer versus mammary epithelial cells. Cancer Research, 52, 6966–6968.
- LIANG, P., AVERBOUKH, L. and PARDEE, A.B., 1993, Distribution & cloning of eukaryotic mRNAs by means of differential display refinements and optimisation. *Nucleic Acids Research*, 21, 3269-3275.
- LIANG, P., BAUER, D., AVERBOUKH, L., WARTHOE, P., ROHRWILD, M., MULLER, H., STRAUSS, M. and PARDEE, A. B., 1995, Analysis of altered gene expression by differential display. Methods in Enzymology, 254, 304-321.
- LINSKENS, M. H., FENG, J., ANDREWS, W. H., ENLOW, B. E., SAATI, S. M., TONKIN, L. A., FUNK, W. D. and VILLEPONTEAU, B., 1995, Cataloging altered gene expression in young and senescent cells using enhanced differential display. *Nucleic Acids Research*, 23, 3244-3251.
- LISITSYN, N., LISITSYN, N. and WIGLER, M., 1993, Cloning the differences between two complex genomes. Science, 259, 946-951.
- LOHMANN, J., SCHICKLE, H. and Bosch, T. C. G., 1995, REN Display, a rapid and efficient method for non-radioactive differential display and mRNA isolation. *Biotechniques*, 18, 200-202.
- LUNNEY, J. K., 1998, Cytokines orchestrating the immune response. Reviews in Science and Techology, 17, 84-94.
- Makowska, J. M., Gibson, G. G. and Bonner, F. W., 1992, Species differences in ciprofibrate-induction of hepaic cytochrome P4504A1 and peroxisome proliferation. *Journal of Biochemical Toxicology*, 7, 183-191.
- MALDARELLI, F., XIANG, C., CHAMOUN, G. and ZEICHNER, S. L., 1998, The expression of the essential nuclear splicing factor SC35 is altered by human immunodeficiency virus infection. Virus Research, 53, 39-51.
- MATHIEU DAUDE, F., CHENG, R., WELSH, J. and McClelland, M., 1996, Screening of differentially amplified cDNA products from RNA arbitrarily primed PCR fingerprints using single strand conformation polymorphism (SSCP) gels. Nucleic Acids Research, 24, 1504-1507.
- MCKENZIE, D. and DRAKE, D., 1997, Identification of differentially expressed gene products with the castaway system. Strategies Newsletter (Stratagene), 10,19-20.
- McClelland, M., Mathieu -Daude, F. and Welsh, J., 1996, RNA fingerprinting and differential display using arbitrarily primed PCR. Trends in Genetics, 11, 242-246.
- MECHLER, B. and RABBITTS, T. H., 1981, Membrane-bound ribosomes of myeloma cells. IV. mRNA complexity of free and membrane-bound polysomes. Journal of Cell Biology, 88, 29-36.
- MEYER, U. A. and ZANGER, U. M., 1997, Molecular mechanisms of genetic polymorphisms of drug metabolism. Annual Review of Pharmacology and Toxicology, 37, 269-296.
- Mohler, K. M. and Butler, L. D., 1991, Quantitation of cytokine mRNA levels utilizing the reverse transcriptase-polymerase chain reaction following primary antigen-specific sensitization in vivo—I. Verification of linearity, reproducibility and specificity. *Molecular Immunology*, 28, 437-447.
- MURPHY, L. D., HERZOG, C. E., RUDICK, J. B., TITO FOIO, A. and BATES, S. E., 1990, Use of the polymerase chain reaction in the quantitation of the mdr-1 gene expression. *Biochemistry*, 29, 10351-10356.
- Nelson, D. R., Koymans, L., Kamataki, T., Stegeman, J. J., Feyereisen, R., Waxman, D. J., Waterman, M. R., Gotoh, O., Coon, M. J., Estabtrook, R. W., Gunsalus, I. C. and Nebert, D. W., 1996, Update on new sequences, gene mapping, accession numbers and nomenclature. *Pharmacogenetics*, 6, 1-42.

- Nishio, Y., Alello, L. P. and King, G. L., 1994, Glucose induced genes in bovine sortic smooth muscle cells identified by mRNA differential display. FASEB Journal, 8, 103-106.
- O'NEILL, M. J. and SINCLAIR, A. H., 1997, Isolation of rare transcripts by representational difference
- ORTON, T. C., ADAM, H. K., BENTLEY, M., HOLLOWAY, B. and Tucker, M. J., 1984, Clobuzarit: species differences in the morphological and biochemical response of the liver following chronic administration. Toxicology and Applied Pharmacology, 73, 138-151.
- PELKONEN', O.; MAENPAA, J., TAAVITSAINEN, P., RAUTIO, A. and RAUNIO, H., 1998, Inhibition and Induction of human cytochrome P450 (CYP) enzymes. Xenobiotica; 28, 1203-1253.
- PHILIPS , S. M., BENDALL , A. J. and RAMSHAW, I. A., 1990, Isolation of genes associated with high metastatic potential in rat mammary adenocarcinomas. Journal of the National Cancer Institute, 82, 199-203.
- PRASHAR, Y. and Weissman, S. M., 1996, Analysis of differential gene expression by display of 3'end restriction fragments of cDNAs. Proceedings of the National Academy of Sciences (USA), 93,
- 659-663.

 RAGNO, S., ESTRADA, I., BUTLER, R. and COLSTON, M. J., 1997, Regulation of macrophage gene expression following invasion by Mycobacterium tuberculosis. Immunology Letters, 57, 143-146.
 - RAMANA, K. V. and Kohli, K. K., 1998, Gene regulation of cytochrome P450—an overview. Indian Journal of Experimental Biology, 36, 437-446.
- RICHARD, L., VELASCO, P. and DETMAR, M., 1998, A simple immunomagnetic protocol for the selective isolation and long-term culture of human dermal microvascular endothelial cells. Experimental
- ROCKETT, J. C., ESDAILE, D. J. and GIBSON, G. G., 1997, Molecular profiling of non-genotoxic hepatocarcinogenesis using differential display reverse transcription-polymerase chain reaction (ddRT-PCR). European Journal of Drug. Metabolism and Pharmacokinetics, 22, 329-333.
- RODRICKS, J. V. and TURNBULL, D., 1987, Inter-species differences in peroxisomes and peroxisome proliferation. Toxicology and Industrial Health, 3, 197-212.
- ROGLER, G., HAUSMANN, M., VOGL, D., ASCHENBRENNER, E., ANDUS, T., FALK, W., ANDREESEN, R., SCHOLMERICH, J. and GROSS, V., 1998, Isolation and phenotypic characterization of colonic macrophages. Clinical and Experimental Immunology, 112, 205-215.
- ROHN, W. M., LEE, Y. J. and BENVENISTE, E. N., 1996, Regulation of class II MHC expression. Critical
- RUDIN, C. M. and THOMPSON, C. B., 1998, B-cell development and maturation. Seminars in Oncology,
- SAKAGUCHI, N., BERGER, C. N. and MELCHERS, F., 1986, Isolation of a cDNA copy of an RNA species expressed in murine pre-B cells. EMBO Journal, 5, 2139-2147.
- SAMBROOK, J., FRITSCH, E. F. and Maniatis, T., 1989, Gel electrophoresis of DNA. In N. Ford, M. Nolan and M. Fergusen (eds), Molecular Cloning—A laboratory manual, 2nd edition (New York: Cold Spring Harbour Laboratory Press), Volume 1, pp. 6-37.
- SARGENT, T. D. and DAWID, I. B., 1983, Differential gene expression in the gastrula of Xenopus laevis.
- SCHENA, M., SHALON, D., HELLER, R., CHAI, A., BROWN., P. O. and DAVIS, R. W., 1996, Parallel human genome analysis: Microarray-based expression monitoring of 1000 genes. Proceedings of the National Academy of Sciences (USA), 93, 10614-10619.
- SCHNEIDER, C., KING, R. M. and PHILIPSON, L., 1988, Genes specifically expressed at growth arrest of
- SCHNEIDER -MAUNOURY, S., GILARDI -HEBENSTREIT, P. and CHARNAY, P., 1998, How to build a vertebrate hindbrain. Lessons from genetics. C R Academy of Science III, 321, 819-834.
- SEMENZA, G. L., 1994, Transcriptional regulation of gene expression: mechanisms and pathophysiology.
- SEWALL, C. H., BELL, D. A., CLARK, G. C., TRITSCHER, A. M., TULLY, D. B., VANDEN HEUVEL, J. and Lucier, G. W., 1995, Induced gene transcription: implications for biomarkers. Clinical
- SINGH, N., AGRAWAL, S. and RASTOGI, A. K., 1997, Infectious diseases and immunity: special reference to major histocompatibility complex. Emerging Infectious Diseases, 3, 41-49.
- SMITH, N. R., LI, A., ALDERSLEY, M., HIGH, A. S., MARKHAM, A. F. and ROBINSON, P. A., 1997, Rapid determination of the complexity of cDNA bands extracted from DDRT-PCR polyacrylamide gels. Nucleic Acids Research, 25, 3552-3554.
- SOMPAYRAC, L., JANE, S., BURN., T. C., TENEN, D. G. and DANNA, K. J., 1995, Overcoming limitations of the mRNA differential display technique. Nucleic Acids Research, 23, 4738-4739.
- ST JOHN, T.P. and DAVIS, R.W., 1979, Isolation of galactose-inducible DNA sequences from Saccharomyces cerevisiae by differential plaque filter hybridisation. Cell, 16, 443-452.
- Sun, Y., Hegamyer, G. and Colburn, N. H., 1994, Molecular cloning of five messenger RNAs differentially expressed in preneoplastic or neoplastic JB6 mouse epidermal cells: one is homologous to human tissue inhibitor of metalloproteinases-3. Cancer Research, 54, 1139-1144.

- SUNG, Y. J. and DENMAN, R. B., 1997, Use of two reverse transcriptases eliminates false-positive results in differential display: Biotechniques, 23, 462-464:
- SUTTON, G., WHITE, O., ADAMS, M. and KERLAVAGE, A., 1995, TIGR Assembler, A new tool for assembling large shotgun sequencing projects. Genome Science and Technology, 1, 9-19.
- Suzuki, Y., Sekiya, T. and Hayashi, K., 1991, Allele-specific polymerase chain reaction: a method for amplification and sequence determination of a single component among a mixture of sequence variants. Analytical Biochemistry, 192, 82-84.
- SYED, V., Gu, W. and HECHT, N. B., 1997, Sertoli cells in culture and mRNA differential display provide a sensitive early warning assay system to detect changes induced by xenobiotics. Journal of
- UITTERLINDEN, A.G., SLAGBOOM, P., KNOOK, D. L. and VUGL, J., 1989, Two-dimensional DNA fingerprinting of human individuals. Proceedings of the National Academy of Sciences (USA), 86,
- ULLMAN, K. S., NORTHROP, J. P., VERWEIJ, C. L. and CRABTREE, G. R., 1990, Transmission of signals from the T lymphocyte antigen receptor to the genes responsible for cell proliferation and immune
- function: the missing link. Annual Review of Immunology, 8, 421-452.

 VASMATZIS, G., ESSAND, M., BRINKMANN, U., LEE, B. and PASTON, I., 1998, Discovery of three genes specifically expressed in human prostate by expressed sequence tag database analysis. Proceedings of the National Academy of Sciences (USA), 95, 300-304.
- VELCULESCU, V.E., ZHANG, L., VOGELSTEIN, B. and KINZLER, K.W., 1995, Serial analysis of gene
- VOELTZ, G. K. and STETZ, J. A., 1998, AuuuA sequences direct mRNA deadenylation uncoupled from decay during Xenopus early development. Molecular and Cell Biology, 18, 7537-7545.
- VOGELSTEIN, B. and KINZLER, K. W., 1993, The multistep nature of cancer. Trends in Genetics, 9,
- WALTER, J., BELFIELD, M., HAMPSON, I. and READ, C., 1997, A novel approach for generating subtractive probes for differential screening by CCLS. Life Science News, 21, 13-14.
- WAN, J. S., SHARP, S. J., POIRIER, G. M.-C., WAGAMAN, P. C., CHAMBERS, J., PYATI, J., HOM, Y.-L., GALINDO, J. E., HUVAR, A., PETERSON, P. A., JACKSON, M. R. and ERLANDER, M. G., 1996, Cloning differentially expressed mRNAs. Nature Biotechnology, 14, 1685-1691.
- WALTER, J., BELFIELD, M., HAMPSON, I. and READ, C., 1997, A novel approach for generating subtractive probes for differential screening by CCLS, Life Science News, 21, 13-14.
- WANG, Z. and BROWN, D. D. 1991, A gene expression screen. Proceedings of the National Academy of
- WAWER, C., RUGGEBERG, H., MEYER, G. and MUYZER, G., 1995, A simple and rapid electrophoresis method to detect sequence variation in PCR-amplified DNA fragments. Nucleic Acids Research,
- WELSH, J., CHADA, K., DALAL, S. S., CHENG, R., RALPH, D. and McClelland, M., 1992, Arbitrarily primed PCR fingerprinting of RNA. Nucleic Acids Research, 20, 4965-4970.
- Wong, H., Anderson, W. D., Cheng, T. and Riabowol, K. T., 1994, Monitoring mRNA expression by polymerase chain reaction: the 'primer-dropping' method. Analytical Biochemistry, 223,
- Wong, K. K. and McClelland, M., 1994, Stress-inducible gene of Salmonella typhimurium identified by arbitrarily primed PCR of RNA. Proceedings of the National Academy of Sciences (USA), 91,
- WYNFORD -THOMAS, D., 1991, Oncogenes and anti-oncogenes; the molecular basis of tumour behaviour.
- XHU, D., CHAN, W. L., LEUNG, B. P., HUANG, F. P., WHEELER, R., PIEDRAFITA, D., ROBINSON, J. H. and Liew, F. Y., 1998, Selective expression of a stable cell surface molecule on type 2 but not type 1 helper T cells. Journal of Experimental Medicine, 187, 787-794.
- YANG, M. and Syrowski, A. J., 1996, Cloning differentially expressed genes by linker capture subtraction. Analytical Biochemistry, 237, 109-114.
- ZHAO, N., HASHIDA, H., TAKAHASHI, N., MISUMI, Y. and SAKAKI, Y., 1995, High-density cDNA filter analysis: a novel approach for large scale quantitative analysis of gene expression. Gene, 156,
- ZHAO, X. J., NEWSOME, J. T. and CIHLAR, R. L., 1998, Up-regulation of two candida albicans genes in the rat model of oral candidiasis detected by differential display. Microbial Pathogenesis, 25, 121-129.
- ZIMMERMANN, C.R., ORR, W.C., LECLERC, R.F., BARNARD, C. and TIMBERLAKE, W.E., 1980, Molecular cloning and selection of genes regulated in Aspergillus development. Cell, 21, 709-715.

Molecular profiling of non-genotoxic hepatocarcinogenesis using differential display reverse transcription-polymerase chain reaction (ddRT-PCR)

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Keywords: ddRT-PCR, non-genotoxic hepatocarcinogenesis, phenobarbital, rat, WY-14,643

SUMMARY

The technique of differential display reverse transcription-polymerase chain reaction (ddRT-PCR) has been used to produce unique profiles of up-regulated and down-regulated gene expression in the liver of male Wistar rats following short term exposure to the non-genotoxic hepatocarcinogens, phenobarbital and WY-14,643. Animals were treated for 3 days, whereupon their livers were extracted and snap frozen. mRNA was prepared from the livers and used for ddRT-PCR. Individual bands from the differential displays were extracted and cloned. False positives were eliminated by dotblot screening and true positives then sequenced and identified.

INTRODUCTION

Safety evaluation of new chemicals usually necessitates the examination of genotoxic and carcinogenic potential using short-term in vitro and in vivo genotoxicity assays augmented by chronic bioassay tests. The short-term assays have proved useful in the early identification of potential genotoxic carcinogens, but their value is limited by observations which suggest that approximately 60% of chemicals identified as carcinogens in life-exposure studies produce mainly negative findings in short-term genotoxicity tests (1,2). Thus, there is currently no reliable and rapid means of evaluating the carcinogenic risk of new chemicals which fall into this latter group of compounds, termed non-genotoxic (or epigenetic) carcinogens.

It is now evident that non-genotoxic carcinogens constitute a group of chemicals which are not only divergent in their interspecies toxicity, but also demonstrate different target organ selectivities and mechanisms of action (3,4). Elucidation of the molecular mechanisms underlying non-genotoxic carcinogenesis is currently underway, but the picture is still far from complete. It is anticipated that a better understanding of the early changes in genetic expression following exposure to non-genotoxic carcinogens will aid development of experimental strategies to identify cellular markers which are diagnostic for this type of toxicity.

Subtractive ddRT-PCR is a recently developed technique which facilitates the preferential amplification of gene products that demonstrate altered expression in target tissue(s) following exposure to chemical stimuli. Furthermore, using this technique, no prior knowledge of the specific genes which are up/down regulated is required. In the current study, we have undertaken to develop a specific and rapid assay for nongenotoxic carcinogens using the technique of ddRT-PCR. This has allowed us to identify characteristic

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patterns of gene regulation following administration of two different non-genotoxic carcinogens (phenobarbital and Wy-14,643) and the subsequent identification of individual gene species which are regulated by this xenobiotic treatment.

MATERIALS AND METHODS

Animals and treatment

Phenobarbital (BDH, Poole, UK; 100 mg/kg/day) or [4-chloro-6-(2,3-xylidino)-2-pyrimidinylthio] acetic acid (Wy-14,643) (Campo, Emmerich; 250 mg/kg/day) was administered by gavage to groups of 3 male Wistar rats (150-200 g) on three consecutive days, whilst control animals received nothing. All animals had free access to food (rat and mouse standard diet, B&K Universal, Hull, UK) and water. The animals were killed on the fourth day, whereupon their livers were excised, sliced into 0.5 cm cubes, snap frozen in liquid nitrogen and then stored at -70°C.

mRNA extraction

Up to 0.25 g of each frozen liver sample was ground under liquid nitrogen using a mortar and pestle. mRNA was extracted from the ground liver using Promega's PolyATtract® System 1000 (Promega, Madison, WI, USA) according to the technical manual. The mRNA was DNase-treated (Promega, final concentration 10 U/ml) before phenol/chloroform extraction and ethanol precipitation. The mRNA was resuspended at a final concentration 500–1000 ng/μl.

ddRT-PCR

This was carried out using the PCR-SelectTM cDNA Subtraction Kit (Clontech, Palo Alto, CA, USA) according to the manufacturer's instructions. Final PCR reactions were run on a 2% Metaphor agarose (FMC, Rockland, MD, USA) gel containing ethidium bromide (Sigma, Dorset, UK) and then overstained for 30 min with SYBR Green I DNA stain (FMC, 1:10 000 dilution in TAE).

Band extraction and cloning

Each discernible band from the differential display pattern was extracted from the gel with a scalpel and the DNA eluted using a Genelute Agarose Spin Column (Supelco, Bellefonte). An aliquot of the eluted DNA (5 µl) was re-amplified using the original ddRT-PCR nested primers and electrophoresed on a 2% agarose gel. The re-amplified band was extracted from the gel (as above) and the eluted DNA ligated directly into the TOPO TA Cloning® vector (Invitrogen, Carlsbad) before transformation in Escherichia coli TOP10F' One Shot[™] cells (Invitrogen).

Stage 1 screening

Twelve transformed (white) colonies from each band were grown up for 6 h in 200 µl LB broth containing ampicillin (Sigma, 50 μ g/ml) and 1 μ l of this amplified by PCR reaction (as specified in ddRT-PCR technical manual). One quarter of the completed reaction was electrophoresed on a standard 2% agarose gel and one quarter on a 2% agarose gel containing HA Yellow (Hanse Analytik GmbH, Bremen, Germany, 1 U/µl) to discern the different cloning products. The remainder was used to prepare duplicate dotblots on Hybond N+ (nylon) membranes (Amersham, Little Chalfont, UK). Cultures containing different cloning products were grown up and a plasmid miniprep prepared from each (Wizard Plus SV Minipreps DNA Purification System, Promega) according to the manufacturer's instructions.

Stage II screening

The duplicate dotblots were probed with: (a) the final differential display reaction; and (b) the 'reverse-subtracted' differential display reaction. To make the 'reverse-subtracted' probe, the subtractive hybridisation step of the ddRT-PCR procedure was carried out using the original tester cDNA as a driver and the driver as a tester. Probing and visualisation were carried out using the ECL Direct Nucleic Acid Labelling and Detection System (Amersham) according to the manufacturer's instructions. Those clones which were positive for (a) but negative for (b), or showed a substantially larger positive signal with (a) compared to (b), were chosen for further analysis.

DNA sequencing

Positive clones as identified above were sequenced on an automated ABI DNA sequencer (Applied Biosystems, Warrington, UK).

B



Fig. 1: (A) Subtractive ddRT-PCR patterns obtained from rat liver following 3-day treatment with WY-14,643 or phenobarbital. Lane 1, 1 kb ladder; lane 2, genes up-regulated following Wy,14-643 treatment; lane 3, genes down-regulated following phenobarbital treatment; lane 4, genes up-regulated following phenobarbital treatment; and lane 6, 1kb ladder. (B) Subtractive ddRT-PCR patterns obtained from rat liver showing relative changes when phenobarbital treated mRNA is subtracted from Wy-14,643-treated mRNA and vice-versa. Lane 1, 1 kb ladder; lane 2, genes showing increased expression following Wy-14,643 treatment compared to phenobarbital treatment; lane 3, genes showing increased expression following phenobarbital treatment compared to Wy-14,643 treatment. See

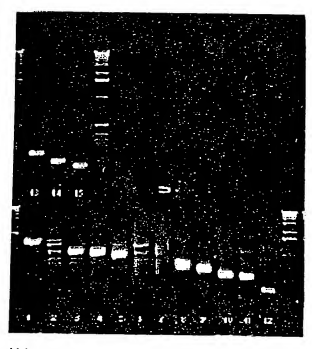


Fig. 2: Re-amplified ddRT-PCR products which were down-regulated following phenobarbital treatment (upregulated bands were also re-amplified but gel not shown). Individual DNA bands excised from gel of ddRTR-PCR reactions were extracted, re-amplified and run on agarose gels to confirm amplification of correct band (numbered). See Materials and Methods for further details.

Table 1: Rat liver genes down-regulated by phenobarbital treatment

Band number (Fig. 2)	Phenobarbital down-regulated		al down-regulated
(Approximate size in bp)	Highestsequ	ence homology	FASTA-EMBL gene identification
1 (1500)		95.3%	Pet mDNA 4 2
2 (1200)		92.3%	Rat mRNA for 3-oxoacyl-CoA thiolase
3 (1000)		91.7%	Rat hemopoxin mRNA
7 (700)	Clone 1	77.2%	R. rattus alpha-2u-globulin mRNA
	Clone 2	94.5%	M. musculus mRNA for Cl inhibitor
	Clone 3	91.0%	Rat electron transfer flavoprotein
8 (650)	Clone 1	86.9%	Mouse topoisomerase 1 (Topo 1) mRNA
	Clone 2	96.2%	Soares 2NbMT M. musculus (EST)
9 (600)	Clone 1	86.9%	Rat alpha-2u-globulin (s-type) mRNA
	Clone 2	82.0%	Soares panies of the soares panies panies of the soares panies of the soares panies of the so
10 (550)		73.8%	Soares panMF19.5 M.musculus (EST)
11 (525)		95.7%	Soares mouse NML M. musculus (EST) NCI_CGAP_Pr1 H. sapiens (EST)
12 (375)		100.0%	R. norvegicus mRNA for ribosomal protein
13 (230)	Cione 1	97 <i>.2</i> %	Soares mouse embryo NbME135 (EST)
	Clone 2	100.0%	Rat fibrinogen B-beta-chain
	Clone 3	100.0%	Rat apolipoprotein E gene
14 (170)		96.0%	Soares p3NMF19.5 M. musculus (EST)
15 (140)		97.3%	Stratagene mouse testis (EST)
Others: (300)		96.7%	R. norvegicus RASP 1 mRNA
(275)		93.1%	Soares mouse mammary gland (EST)

EST = expressed sequence tag.

Bands 4-6 were shown to be false positives by dotblot analysis and, therefore, not sequenced.

Table II: Rat liver genes up-regulated by phenobarbital treatment

Band number Approximate size in bp)	Phenobarbital up-regulated		
	Highestseque	ence homology	FASTA-EMBL gene identification
5 (1300)		93.5%	Dat marks
7 (1000)		95.1%	Rat cytochrome P450IIB1 mRNA for rat preproalbumin
			Rat serum albumin mRNA
8 (950)		98.3%	NCI_CGAP_Pr1 H. sapiens (EST)
10 (850)	•	95.7%	Rat cytochrome P450IIB1
11 (800)	Clone 1	94.9%	Rat cytochrome P450IIB1
12 (750)	Clone 2	75.3%	Rat cytochrome p450-L (p450IIB2)
		93.8%	Rat TRPM-2 mRNA
15 (600)			Rat mRNA for sulfated glycoprotein
		92.9%	mRNA for rat preproalburnin
16 (550)			Rat serum albumin mRNA
	Clone 1	95 <i>.</i> 2%	Rat cytochrome P450IIB1
21 (350)	Clone 2	93.6%	Rat haptoglobulin mRNA partial alpha
		99.3%	R. norvegicus genes for 18S, 5.8S & 28S rRNA

EST = expressed sequence tag.

Identification of differentially-regulated genes

Gene-sequences were identified using the FASTA programme (http://www.ebi.ac.uk/htbin/fasta.py?request) to search all EMBL databases for matching DNA sequences.

RESULTS

Figure 1A,B shows the ddRT-PCR patterns of genes showing altered expression in rat liver following 3 day treatment with phenobarbital or Wy-14,643. Individual bands were isolated from the phenobarbital-modulated patterns (both up- and down-regulated), re-amplified (Fig. 2), cloned, screened for false positives and then identified. Those xenobiotic-modulated gene products identified to date are listed in Tables I and II.

DISCUSSION

The advent of combinatorial chemistry has led to the synthesis of millions of new chemical compounds, many of which may be potentially useful in pharmaceutical, agricultural or industrial applications. However, whilst there are tests available for those posing a genotoxic activity, there remains no short-term assay able to identify those chemicals which may belong to the non-genotoxic group of carcinogens.

We have used an adaptation of the subtractive hybridisation method – ddRT-PCR – to produce characteristic profiles or 'fingerprints' of those genes which are up-regulated or down-regulated in male rat liver following acute exposure to test chemicals. The ddRT-PCR profiles are characteristic and unique for each of the 2 compounds studied to date.

A number of those gene species showing altered expression following phenobarbital treatment have been cloned and identified (Tables I & II). It is interesting to note the presence of CYP2B2 in the up-regulated genes. This would, of course, be expected following exposure to phenobarbital and serves as a positive control for the method. Other genes which one might normally expect to be up-regulated do not appear in the table. However, it should be noted that not

all bands seen on the differential display were extracted and re-amplified due to their being too faint or too close to other bands to accurately excise. Furthermore, it has been well documented [(5) and references therein] that a single band extracted from a differential display often represents a composite of heterogeneous products. We are currently examining new methods to:

(i) improve resolution of the differential display patterns (including 2-D agarose gels); and (ii) distinguish those ddRT-PCR products which are identical in size, but different in sequence.

Our future efforts will be directed towards determining the extent of modulation of a number of the genes reported herein using semi-quantitative RT-PCR. This should reveal the extent of changes in expression of key gene products which may be involved in non-genotoxic hepatocarcinogenesis and thus help increase understanding of this process. Furthermore, it is anticipated that aligning ddRT-PCR profiles of different non-genotoxic agents found in responsive and non-responsive species may enable identification of those genes which are mechanistically relevant to the non-genotoxic hepatocarcinogenic process. Accordingly, this approach lends itself well to the identification, characterisation and sub-classification of possible different classes of non-genotoxic carcinogens.

ACKNOWLEDGEMENT

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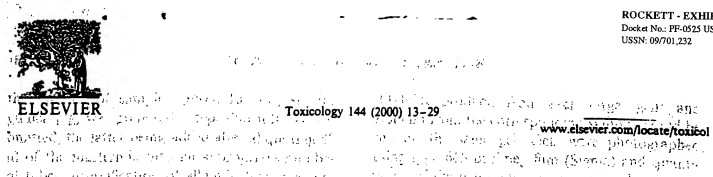
REFERENCES

- Parodi S. (1992): Non-genotoxic factors in the carcinogenic process: problems of detection and hazard evaluation. Toxicol. Lett., 64/65, 621-630.
- Ashby J. (1992): Prediction of non-genotoxic carcinogenesis. Toxicol. Lett., 64/65, 605-612.
- Grasso G. and Sharratt M. (1991): Role of persistent, non-genotoxic tissue damage in rodent cancer and relevance to humans. Annu. Rev. Pharmacol. Toxicol., 31, 253-287.
- Lake B. (1995): Mechanisms of hepatocarcinogenicity of peroxisome-proliferating drugs and chemicals. Annu. Rev. Pharmacol. Toxicol., 35, 483-507.
- Smith N.R., Li A., Aldersley M., High A.S., Markham A.E., Robinson P.A. (1997): Rapid determination of the complexity of cDNA bands extracted from DDRT-PCR polyacrylamide gels. Nucleic Acids Research 25 (17), 3552-3554.

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Use of suppression-PCR subtractive hybridisation to identify genes that demonstrate altered expression in male rat and guinea pig livers following exposure to Wy-14,643, a peroxisome proliferator and non-genotoxic hepatocarcinogen

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Abstract

Understanding the genetic profile of a cell at all stages of normal and carcinogenic development should provide an essential aid to developing new strategies for the prevention, early detection, diagnosis and treatment of cancers. We have attempted to identify some of the genes that may be involved in peroxisome-proliferator (PP)-induced non-genotoxic hepatocarcinogenesis using suppression PCR subtractive hybridisation (SSH). Wistar rats (male) were chosen as a representative susceptible species and Duncan-Hartley guinea pigs (male) as a resistant species to the hepatocarcinogenic effects of the PP, [4-chloro-6-(2,3-xylidino)-2-pyrimidinylthio] acetic acid (Wy-14,643). In each case, groups of four test animals were administered a single dose of Wy-14,643 (250 mg/kg per day in corn oil) by gastric intubation for 3 consecutive days. The control animals received corn oil only. On the fourth day the animals were killed and liver mRNA extracted. SSH was carried out using mRNA extracted from the rat and guinea pig livers, and used to isolate genes that were up and downregulated following Wy-14,643 treatment. These genes included some predictable (and hence positive control) species such as CYP4A1 and CYP2C11 (upregulated and downregulated in rat liver, respectively). Several genes that may be implicated in hepatocarcinogenesis have also been identified, as have some unidentified species. This work thus provides a starting point for developing a molecular profile of the early effects of a non-genotoxic carcinogen in sensitive and resistant species that could ultimately lead to a short-term assay for this type of toxicity. © 2000 Elsevier Science Ireland Ltd. All rights reserved.

Keywords: Wy-14,643; Peroxisome proliferator; Non-genotoxic hepatocarcinogenesis; Suppression PCR subtractive hybridisation; RT-PCR; Rat; Guinea pig; Gene regulation; Differential gene display; Gene profiling

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Introduction

The advent of combinatorial chemistry and imputer-aided drug design has led to a recent osurge in the number of chemical compounds at have potential therapeutic, agricultural and dustrial applications. Although it has been sugested that the contribution of synthetic chemicals the overall incidence of human cancer is low, ere still remains an absolute requirement to aluate all new chemicals for toxic and carcinonic potential. The latter is one of the most oblematic areas of chemical safety evaluation id is usually carried out using short-term in vitro id in vivo genotoxicity assays augmented by ronic bioassay tests. The short-term assays have oved useful in the early identification of poten-I genotoxic carcinogens, but their value is limobservations that suggest that proximately 60% of chemicals identified as carlogens in life-exposure studies produce mainly gative findings in short-term genotoxcity tests shby, 1992; Parodi, 1992). Thus, there is curitly no reliable and rapid means of evaluating carcinogenic risk of new chemicals that fall o this latter group of compounds, termed nonnotoxic (or epigenetic) carcinogens.

One approach to addressing this problem is to cidate the molecular mechanisms by which own non-genotoxic carcinogens act. It should n be possible to identify common factors/ chanisms that can serve as early biomarkers of cinogenic potential for new chemicals. To this l, a large number of groups have reported on various effects of non-genotoxic compounds various animal species (Marsman et al., 1988; ce et al., 1993; Cattley et al., 1994; Hayashi et 1994; Human and Experimental Toxicology, 4; Anderson et al., 1996). However, the mechstic picture is still far from complete with many those genes involved in the carcinogenic proremaining unknown, and their identification efore remains a key goal in elucidating the ecular mechanisms by which non-genotoxic inogenesis occurs.

ubtractive hybridisation (SH) and related techogies such as representational difference analy-(RDA) (Hubank and Schatz, 1994) and

out Aries as agreement Dominion spring a differential display (DD) (Liang and Pardee, 1992) can be used to aid the isolation of genes of the showing altered expression in target tissues following exposure to a chemical stimulus. These $\frac{1}{2}$ techniques can also be used to identify differential gene expression in neoplastic and normal cells, (Liang et al., 1992), infected and normal cells (Duguid and Dinauer, 1990), differentiated and theres. undifferentiated cells (Sargent and Dawid, 1983; Guimaraes et al., 1995), activated and dormant cells (Gurskaya et al., 1996; Wan et al., 1996), different cell types (Hedrick et al., 1984; Davis et al., 1984) amongst others. Most importantly, using such approaches, no prior knowledge of the specific genes that are upregulated/downregulated is required.

Using a variation of SH, termed suppression-PCR subtractive hybridisation (SSH) (Diatchenko. et al., 1996), we have previously reported the isolation of a number of genes showing altered expression in male rat liver following acute exposure to phenobarbital (Rockett et al., 1997). In the current work we have used the same experimental approach to isolate genes that are differentially expressed in the livers of male rats and guinea pigs following short-term (3-day) exposure to the peroxisome proliferator (PP) and nongenotoxic hepatocarcinogen, Wy-14,643. We have isolated and identified a number of gene species, some of which may be important in the induction or · protection against, non-genotoxic hepatocarcinogenesis.

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2. Materials and methods

2.1. Animals and treatment

All animal experiments were undertaken in accordance with Her Majesty's Home Office Department guidelines under the auspices of approved personal and project licences. Male Wistar rats (150-200 g) and male Duncan-Hartley guinea pigs (250-300 g) were obtained from Kingman and Bantam (Hull, UK). Upon receipt, both groups were randomly assigned into two groups of four. They were maintained on a rat, mouse or guinea pig standard diet (B&K Univer-

sal, Hull) and a daily cycle of alternating 12-h Metaphor agarose gel (FMC, Rockland, USA) . periods of dark and light. The room temperature was maintained at 19°C and a relative humidity of 55%. The animals were acclimatised to this envi-.. ronment for 7 days before treatment commenced. [4-chloro-6-(2,3-xylidino)-2-pyrimidinylthio] acetic acid (Wy-14,643; Campo, Emmerich; 250 mg/kg per day in corn oil) was administered by gavage to the treated groups of rats and guinea pigs on 3 consecutive days, whilst control groups received an equal volume of corn oil only. During this time, all animals had free access to food and water. The animals were killed by cervical dislocation on the fourth day, and their livers immediately excised, weighed; sliced into approximately 0.5-cm cubes, snap frozen in liquid nitrogen and stored at -70°C.

2.2. mRNA extraction

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Approximately 0.25 g of each frozen liver sample was ground under liquid nitrogen using a mortar and pestle. Messenger RNAswas extracted from the ground liver using the PolyATtract® System 1000 kit (Promega, Madison, USA) according to the technical manual provided by the manufacturers. The mRNA was DNase-treated (RQ Rnase-free Dnase, Promega, final concentration 10 U/ml) before phenol/chloroform extraction and ethanol precipitation. The mRNA was redissolved at a final concentration 500-1000 ng/ $\mu l.$

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2.3. cDNA Subtraction

This was carried out using the PCR-SelectTM cDNA Subtraction Kit (Clontech, Palo Alto, USA) according to the manufacturer's instructions. Subtractions were carried out with mRNAs derived from single animals. The mRNA from the remaining three animals in each group was later used for quantitative RT-PCR analysis of specific genes.

2.4. Band extraction and cloning

The secondary PCR reactions from the cDNA subtraction procedure were run on a 2%

containing 0.5 µg/ml ethidium bromide (Sigma, Dorset, UK). One times TAE (0.04 M Tris-acetate, 0.001 M EDTA) was used to prepare the gel and as the running buffer. After running for 6-7 h at 3.75 V/cm, the gel was overstained for 30 min with SYBR Green I DNA stain (FMC, 1:10 000 dilution in 1 × TAE). Each discernible band of the differential display pattern was extracted from the gel with a scalpel and the DNA eluted using a GeneluteTM agarose spin column (Supelco, Bellefonte, (USA). Five microlitres of the eluted DNA was reamplified using the original nested (secondary) PCR primers supplied with the PCR-Select™ cDNA subtraction kit. The PCR products were electrophoresed on a 2% standard agarose gel (Boehringer Mannheim, East Sussex, UK) and ethe reamplified target bands extracted from the gel as above. The eluted DNA was immediately ligated into a TOPO TA Cloning® vector (Invitrogen, Carlsbad, USA) before transformation in Escherichia coli TOP10F' One Shot™ (Invitrogen).

2.5. Colony screening

2.5.1. Stage I

Eight transformed (white) colonies from each band were grown up for 6 h in 200 µl LB broth containing ampicillin (Sigma, 50 mg/ml). One microlitre of this was subjected to PCR using the same conditions and nested primers as described above. One tenth (2 µl) of the completed PCR reaction was electrophoresed on a 2% standard agarose gel and one tenth on a 2% standard agarose gel containing HA red (Hanse Analytik GmbH, Bremen, Germany, 1 U/ml) to discern the differentially cloned products. The remainder of the PCR reaction was used to prepare duplicate dotblots on Hybond N+ membranes (Amersham, Little Chalfont, UK).

2.5.2. Stage 11

The duplicate dotblots were probed with (a) the final differential display reaction and (b) the 'reverse-subtracted' differential display reaction. To make the 'reverse-subtracted' probe, the subtractive hybridisation step of the differential display the state of the state of the state of

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RT-PCR procedure was carried out using the original tester (treated) mRNA as the driver and he original driver (control) mRNA as the tester. Probing and visualisation were carried out using he ECL direct nucleic acid labelling and detection system (Amersham, Little Chalfont, UK) according to the manufacturer's instructions. Those lones that were positive for (a) but negative for b), or showed a substantially larger positive signal with (a) compared to (b), were selected for DNA sequence analysis.

2.6. DNA sequencing

The remainder of the cultures (prepared in tage I screening) containing different cloning roducts (as discerned in the two screening steps) were grown up overnight in 5 ml LB broth containing ampicillin (50 mg/ml). A plasmid miniprep as prepared from each (Wizard Plus SV linipreps DNA purification system, Promega) ccording to the manufacturer's instructions. The loned inserts were sequenced on an automated BI DNA sequencer (Applied Biosystems, Warngton, UK) using the M13 forward primer GTAAAACGACGGCCAGT) or M13 reverse rimer (AACAGCTATGACCATG).

7. Identification of differentially regulated genes

Gene sequences thus obtained were identified sing the FASTA 3.0 programme (Lipman and earson, 1985; Pearson and Lipman, 1988) (http://www.ddbj.nig.ac.jp/E-mail/homology.html) to arch all EMBL databases for matching DNA quences. Each clone sequence was submitted in e forward and reverse direction, and the one turning the highest statistical probability of atch to a known sequence was noted. Sequence omologies between our submitted clone sequence id the queried database sequence were deterined (by FASTA) over a region of at least 60 se pairs.

3. RT-PCR analysis of selected candidate genes

cDNA sequences of the target genes were obned from the NIH gene database (GenBank at

http://www.ncbi.nlm.nih.gov/Web/Search/index. html) and the computer programme GENES JOCKEY (BioSoft, Cambridge, UK) used to select primer pairs from these sequences. Where guinea pig sequences were available, rat and guinea pig sequences were aligned and primers chosen from regions of homology. If guinea pig sequences were not available, rat and human sequences were used. In cases where exact homology could not be found, the sequence from the rat was used. In the case of CD81 only, no rat or guinea pig sequences were available and so mouse and human sequences were aligned and a primer pair chosen from a region of homology. Primers (obtained from Gibco-BRL, Paisley, UK) were dissolved at a concentration of 50 pmol/µl in sterile distilled water and stored at -20° C. The primer pairs used plus other reaction parameters are shown in Table 1. mRNA was extracted (as described above) from all four treated animals and from three animals in the control group. Integrity of the eluted mRNA was confirmed on a 2% agarose gel, and the concentration and purity were measured using a Genequant II spectrophotometer (LKB, Bromma, Sweden) and then diluted to 10 ng/µl. One microlitre of this latter solution was... used per RT-PCR reaction.

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RT-PCR was carried out in a single tube (50 µl) reaction using the Access RT-PCR system (Promega) according to manufacturer's instructions. In the kinetic and quantitative analyses, omission of RNA was used as a control for the presence of any contaminating DNA. After obtaining a PCR signal of the correct size and optimising the reaction conditions, each PCR product was digested with between two and four separate restriction enzymes. Specific restriction patterns were thus obtained, which further confirmed the identity of the PCR products as being the original target genes. Kinetic analysis (14–32 cycles) was then performed in each case to determine the location of the mid-log phase.

For the semi-quantitative analysis of each target gene, RT-PCR reactions were carried out in triplicate for each sample to reduce the effect of intertube RT-reaction variations (Kolls et al., 1993) and pipetting errors. For each gene, a mastermix containing enough reagents for three times

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100698 (rat) TGGAGAGA CTTAG G6/59	i ranscript	Genbank ac- cession No.	Primer sequences	.,	Size of rat PCR product (bp)		perature		No. of rat/guin	PCR c	ycle
100698 (rat) TGGAGAGA				downstream			eserci.		n T		
CANDED C	Albumin	J00698 (rat)	TGGAGAGA-	CTTAG.	436				8. 1. 	P.(
CACCAGC CACC	Bifunctional enzyme	K03249 (rat)	GAGC. CTTCAAAGC GCACC. CACTTCTTCT.	CAAGTCTCAGC CA TGGCAATGATG	G 347		्रक्ति इ.स्तु दिश	er bed merga	15/22 21/-2-112	n Naday On Jawa	
M14972 (rat) CTGAGG GATTGT 357 57/4-1 196	CYP2C11	J02657 (rat)	CACCAGC CCATCATGACC.	GAAGTCCCGAG	410		s Art Er Fratt	an si Mi d	unverg E P	alas b nesad	
CATGAG GATCTGATC	CYP4A!	M14972 (rat)	CTGAGG GATGGCTGCAC-	GATTGT GGCCTTTG-	357		*5.3 193	ociji Na	5 5 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	i vi. Fires	•
TAATGG	Catalase	M11670 (rat)	CATGAG ACCAAATACTC. CAAGGCAAAGG	GATCTGATC GCCCTG. GTCAGTCTTG.	<u>5</u> 450	-	٠	is G	27/- 55 27/-	ei in	
Minuse GCTGGCTGG GAACTGCTTCA RNCCP23 GATCTGCTGGTTG 341 50/- 70 20/- 70 20/- 70 70/- 70 70/-	CD81 (TAPA-1)	X59047	ATTTCGTCTTCTG		· · · · · · · · · · · · · · · · · · ·		fjer t	our.	्र		•
CGCACCAT	Contrapsin-like	(mouse) RNCCP23	GCTGGCTGG GACTATGTGAG.	GAACTGCTTCA GTCTCTGGTTG	34I		rin	Strate i Barrier	1117.ks	10000	•
D38380 (rat) AGCTGTGT- GAGGAGACC- 360 57/59 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5	Parathymosin-a (Zn ²⁺ binding profein)	X64053 (rat)	CGGCACCAT.	CAAGCI TTGTGTGTTCCT	382		MM (C	udiliri Villar	Sper €		
GTCCAGG GAA GGAT U00273 (rat) GGAT GCAGTTTCAGCT OF CASS Solve Solve Solve CAGGCAGG ANA CAAGGCA CAAGGCA GAGTTCCAGG GAGTTCCAGG GAGTTCCAGG GAGTTCCAGG GAGTTCCAGG GAGTTCCAGG GAATAACA GAATTCCAGG GAAGCAGTG GAAGCAGG GAAGCAGTG G	Transferrin	D38380 (rat)	AGCTGTGT.	GAGGAGAGCC.	360	ر و	**************************************	dion.	22/22		-
U/a CGACGTTTC- TGTTGCGGCA- 318 55/- C CAAGGCA GAGTGGA CAAGGCA GAGTGGA D21058 (rat) CAAATAACA- CTCCATCCAGG GAAGCAGTG- CTCCATCCAGG GAAGCAGTG- CTCCATCCAGG GAGC	UDP-GT		ACCA 3AT-	AGITIG- ITCAGC IGCT	MARIN (1997) (1997) (1997) (1997) (1997)	20/-4/2	decided	i Told Dol Ngjarjere	(31.04), miqu a		
D21058 (rat) CAAATAACA- GACTTCCAG- 4333 R S R S R S R S R S R S R S R S R S	Down Unknown-1	ท/ล	CGACGTTTC	TGTTGCGGCA.	318	`\	agair	r Zuci Milita		à Gri	
	Znazglycoprotéin	D21058 (rat)		ي	9 - 9: 88 60	augungs	તા એક જ	South or		:1, [992]	

III Barlan in plant transfer to the setting of the

the number of samples (seven for rat, six for guinea pig) was prepared except that mRNA was omitted, the latter being added after aliquoting 49 ul of the mastermix into an appropriate number of tubes. Amplification of albumin (the reference gene) was carried out in separate tubes since the mid-log phase of this gene is at a much lower cycle number than the target genes due to its high abundance. All RT-PCR products were analysed on 2% agarose gels containing 0.5 μg/ml ethidium bromide. The target gene samples were loaded on the gel first and run in at 3 V/cm for 10 min. The corresponding albumin samples were then loaded and the gel run for a further 1/2 h. In this way, all

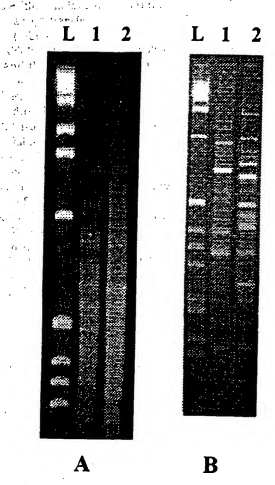


Fig. 1. Final displays of differentially expressed genes that were (1) upregulated and (2) downregulated in rat (A) and guinea pig (B) livers following 3-day treatment with Wy-14,643. mRNA extracted from control and treated livers was ised to generate the differential displays using the PCR-Select :DNA subtraction kit (Clontech). Lane (L) is a 1 Kb DNA Ladder standard and 10 µl of secondary PCR reaction were oaded in all other lanes.

RT-PCR products from each target gene and albumin from the corresponding samples could be run on the same gel. Gels were photographed using type 665 posi-neg film (Sigma) and quantitation of the band intensity was carried out using a dual wavelength flying spot laser scanner densitometer (Shimadzu). • (**

2.9. Statistical analysis 🚟

Statistical analysis of unpaired samples was carried out using the two-tailed Student's t-test. Values were considered statistically significant at P < 0.05 or less.

in, 273 3. Results

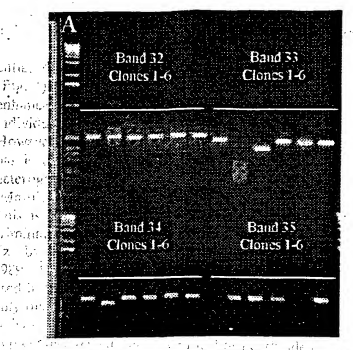
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3.1: Cloning and screening of transcripts

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For both the rat and guinea pig experimental groups, cDNA subtraction was carried out in the forward (control driving tester) and reverse (tester driving control) directions to isolate both upregulated and downregulated mRNA species respectively. Using a standard primary hybridisation time of 8 h we obtained a substantial amount of non-specific products in all the final differential displays (data not shown). This background smearing was almost completely removed by reducing the primary hybridisation time to 4 h (CLONTECHniques, 1996). Fig. 1 shows theddRT-PCR patterns of genes showing altered expression in rat and guinea pig liver following 3-day treatment with Wy-14,643. The profiles are unique for each species, and in each case the profile for the upregulated genes (control mRNA driving tester mRNA) is different to that obtained for the downregulated genes (tester mRNA driving control mRNA).

The practical outcome of the SSH method is that a series of differentially expressed genes is observed as a ladder on an agarose gel. The majority of these gene fragments fall within the 150-2000 bp range, with bands up to 5 Kbp occasionally being observed. Each band may theoretically consist of one or more products of similar size, as the gel has a maximum resolution



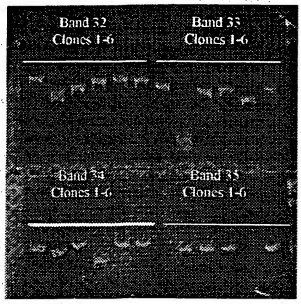


Fig. 2. Discrimination of different ddRT-PCR products having the same molecular size using HA-red. Gel (A) is a 2% standard agarose gel. Gel (B) is a 2% standard agarose gel containing 1 U/ml HA-red. Band numbers refer to the sequential bands (largest to smallest) extracted from the original display of genes upregulated in rat liver following 3-day treatment with Wy-14,643. Ten micorlitres of each PCR reaction were loaded per lane.

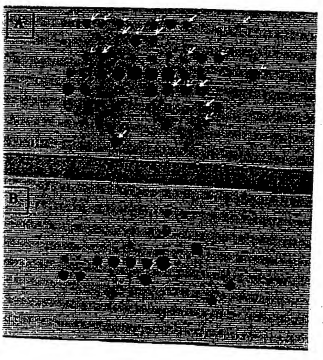
of approximately 1.5% (3 bp per 200). In addition, there may be two or more products that are the same size, but have a different sequence.

Therefore some form of discrimination must be employed to isolate as many of these products as possible. HA-red screening (Geisinger et al., 1997) of a number of clones derived from each band provided a means to discriminate between different gene species of the same size. A typical example of such a gel is shown in Fig. 2. In total, 88 and 48 apparently different clones were obtained from the final differential expression patterns of upregulated and downregulated rat genes, respectively. Sixty nine and 89 apparently different clones were obtained from the final differential expression patterns of the upregulated and downregulated guinea pig genes, respectively.

Having identified as many different candidate gene products as possible in the screening step I, a second screening step was carried out on every clone to confirm those that represented true differentially expressed genes. This is necessary since no subtraction technique is 100% efficient. The approach we used, termed PCR-select differential screening (as recommended in Clontech's PCR-select cDNA subtraction kit protocol), utilises the forward and reverse subtractions as an aid to screening for the true differentially expressed genes (CLONTECHniques, 1997). Because these probes have already undergone subtraction, they have been enriched for differentially expressed genes and are therefore more sensitive than unsubtracted driver/tester cDNA probes for detecting true differential expression. All the clones that were isolated from each display were dotblotted and probed with the display from which they was obtained, plus the corresponding reverse-subtracted display. An example of such a blot is shown in Fig. 3. Clones corresponding to authentic differentially expressed mRNAs hybridised with the subtracted cDNA probe, but not the reverse-subtracted probe. We also included in the authentic positives, those clones that gave a substantially greater signal with the subtracted probe compared to the reverse-subtracted probe. False positives hybridised with either both probes or with neither probe. Of the original 88 upregulated and 48 downregulated rat clones selected for this screening step, 28 (32%) and 15 (31%) respectively, were found to be true positives. In the rat. 8 (100%) of the true positive upregulated genes Table 2) and 11 (73%) of the true positive downegulated genes (Table 3) were non-redundant. Of the original 69 upregulated and 89 downregulated uinea pig clones selected for this screening step, 8 (70%) and 37 (42%) respectively, were found to e true positives. Thirty six (75%) of the upregulated genes (Table 4) and 33 (89%) of the downgulated genes (Table 5) were non-redundant.

2. Identification of clones

On sequence analysis it was found that some ones were unsequencable in the first instance 113 forward primer) due to long polyA runs at appeared to prematurely terminate the senencing reaction. These clones were therefore sequenced from the opposite direction using the 13 reverse primer. Those xenobiotic-modulated ne products identified to date are listed in Taes 2 and 3 (rat) and Tables 4 and 5 (guinea pig).



3. Dot blots of clones of putative upregulated gene species ted from guinea pig liver following 3-day treatment with 14,643. All clones identified in the stage I screening step methods) were blotted and probed with (A) the differendisplay from which they originated (control driving ed) and (B) the reverse subtraction (treated driving con-Arrows indicate some of the true differentially expressed

Table 2
Identification of genes that were upregulated in male rat liver following 3-day treatment with WY-14.643.

FASTA-EMBL gene identification (rat unless otherwise stated)		Sequence homology* (%)
Carnitine octanoyl transferase		99
NCI_CGAP_Lil (H. sapiens) (EST ^b)	. HS1275949	98
Peroxisomal enoyl hydratase-like	RN08976	98
protein		
Liver fatty acid bind- ing protein	V01235	96
Soares mouse p3NMF19.5 M.	AA038051	96
musculus cDNA clone		-
Cytochrome p450IVA1	RNCYPLA	94
Mit. 3-hydroxyl-3- methylglutaryl CoA synthase	RNHMGCOA	94
Rabgeranylgeranyl transferase compo-	RNRABGERA	94
nent B Genes for 18S, 5.8S, and 28S ribosomal RNAs	RNRRNA	94
Carnitine acetyl transferase (mouse)	MMRNACAR	92
Soares mouse NML (EST)	MM1157113	92
Bone marrow stromal fibroblast (H. sapi-	AA545726	92
ens) cDNA clone HBMSF2E4 (EST)	y	
7.5dpc embryo (mouse) (EST)	AA408192	92
Alpha-1-macroglobu- lin	RNALPHIM	91
Transferrin	RNTRANSA	91
Lecithin:cholesterol acyltransferase	RNU62803	90
Zn-α2-glycoprotein	RNZA2GA	90
Serum albumin	RNJALBM	89
phate 1-phospho- hydrolase	RNFBP	88
oares mouse melanoma (EST) (S ^c)	AA124706	88
oares mouse 3NbMS (EST) (AS ^c)	AA154039	88

(B) (0)

35

Table 2 (Continued)

FASTA-EMBL gene identification (rat unless otherwise stated)		hamala — # (0/)
17-β-hydroxsteroid de- hydrogenase (17) Soares mouse	RN17BHDT2 (*)	or the these
p3NMF19.5 (EST) Peroxisomal enoyl- CoA:hydratase -3-	RNPECÓA	385 141 °C 1, 12.11
hydroxyacyl CoA bifunctional enzyme Integral membrane	The State of State of the State	The state of the s
protein, TAPA-1 (CD81) (mouse) Soares mouse lymph	gone to Milita	
node (EST) H. sapiens (clone	L40401	7 6 an in an
zap128) mRNA Lysophospholipase ho-	HSU67963	76
mologue (human) Soares mouse lymph node (EST)	AA217044	

a Refers to the nucleotide sequence homology between the cloned band isolated from the differential display and the corresponding gene derived from the EMBL gene sequence bank.

In all cases, both the forward and reverse sequence of the target clones were analysed and the gene having the highest statistical homology noted.

3.3. RT-PCR analysis of selected clones

The results of a typical RT-PCR semi-quantitation experiment for transferrin in the rat is given in Fig. 4 and the results for a total of 12 selected genes in both the rat and guinea pig are shown in Table 6.

Table 3 Identification of genes that were downregulated in male rat liver following 3-day treatment with Wy-14,643

identification (rat un- less otherwise stated)	Accession No. Sequence des should be
NCI_CGAP_Lil (H.	AA484528 99
sapiens) (ESTb)(Sc)	of the Briggs which to Intife whom:
	AA469320 Shine 99 to Review
sapiens) (EST)(ASc)	and the letter of the control of the
UDP-glucuronosyl-	RN06273 98
transferase	The state of the s
(UGT2B12)	summer the second
Complement compo-	RNC3
nent c3	化氯化甲基甲基苯甲基甲基甲基甲基
Soares mouse pla- 🗀 🗄	AA023305 m. 5 . 96 . 5 . 5 . 6 . 6
centa (S)	street along the contract of
whe (criminhausee) 509'	PTRGMC 96
rRNA (AS)	The area of the first autilities
Rat CYP2C11	RNCYPMI 95
Ribosomal protein S5	*RNRPS5" 10" at 94" (122) [123]
Transthyretin :	RNTTHY Hill 194 Control
Contrapsin-like	RNCCP23
protease inhibitor.	
Prostaglandin F2a (S)	RN26663 84
B-2-microglobulin	RNB2MR 84 Fig. 1
(AS)	等的是有一种的一种。 化二十二十二十二十二十二十二十二十二十二十二十二十二十二十二十二十二十二十二十
Apolipoprotein C-III	RNAPOA02-1 C825FFGF EX
Parathymosin-alpha (zinc2+-binding protein)	RNIIZNBP 75

a Refers to the nucleotide sequence homology between the cloned band isolated from the differential display and the corresponding gene derived from the EMBL gene sequence bank.

4. Discussion

It is now apparent that all cancers arise from accumulated genetic changes within the cell. Although documenting and explaining these changes presents a formidable obstacle to understanding the different mechanisms of carcinogenesis, the experimental methodology is now available to begin attempting this difficult challenge. In order to begin the elucidation of the molecular mechanisms involved in non-genotoxic hepatocarcino-

EST is 'expressed sequence tag' — a gene of as yet unknown identity and function.

^c Where sequence homologies were equal in both directions of the isolated band, both the sense (S) and antisense (A) identities are given.

^b EST is 'expressed sequence tag' — a gene of as yet unknown identity and function.

^c Where sequence homologies were equal in both directions, both the sense (S) and antisense (A) identities are given.

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remainer Contract of the 1994 in the contract enesis, we have used SSH to identify a number of 100. Lake et al., 1989; Makowska et al., 1992; Lake et 1 enes that are upregulated or downregulated in the nale_rat_and_guinea pig_livers following short. erm, exposure to the PP, Wy-14,643. We have sed the rat model to represent a species susceptile to the non-genotoxic carcinogenic effect of Ps and the guinea pig as a resistant species

Orton et al., 1984; Rödricks and Turnbull, 1987;

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1906) Burtham to the out.

三边运输 新国的统治的 (through et no 1907) (Kra

mothers and drokes are brainfus or emotioning to al., 1993). They from the terms I so the employeen

Gurskaya et al. (1996), who originally developed the SSH technique, cloned the products of the the secondary PCR reaction and screened a small at number of randomly selected colonies for differentially expressed clones using northern hybridisation. However, we decided against this approach ! the chardshor of specific and school to become be

er, van Mero i gynt of vrinces, haver est instientification of genes that were upregulated in male guinea pig liver following 3-day treatment with WY-14.643

		1 1 2 2 2 2 2 2	15 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
ASTA-EMBL gene identification (guinea pig unless otherwise stated)	receis mai j	Accession 1	No. 13. Sequence 19 the Care
State a room to the to the first the same	i en etterrie ei	The Alexander	homology" (%)
arboxylesterase		AB010634	97
omplement C3 protein (GPC3)	sin's gur, chy	M34054	THE SHOT SHOT SHOT
ytosolic aldehyde dehydrogenase (sheep)	Sacra dia dia	T112761	1 92 c L + h a n a n (2) e
italase (human)	The state of the s	X04076 :	180 6 89 3 4 5 6 6 6 6 6 6 6 6 6
itochondrial aspartate aminotransferase (pig)		M11732	
ongation factor-1-alpha (rabbit)	- 1 / 2 1 4 4 22 2 4 1 4 2 2 2 4 4 2 2 2 2 4 4 2 2 2 2 2 2 2	X62245	
ongation factor-1-alpha (rabbit) CI_CGAP_Br2 H. sapiens cDNA clone (EST) (Similar to chick mit.	phosphoenolpyru-	AA587436	is th <mark>88</mark> known etches
vate carboxykinase)		1111507450	on said the same there of
lpha-1-antiproteinase S		M 57270	12. 183 1 14. 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
-formyltetrahydrofolate dehydrogenase (rat)			183
bosomal protein L6 (rat)		X87107	83.
ares pregnant uterus Nb (EST) (mouse)		AA156847	83
itochondrial citrate transport protein (human)	10.0	L77567	80
toplasmic chaperonin hTRiC5 (human)		U17104	80
pha-1-antiproteinase F		M57271	77
eterogeneous nuclear ribonuclearprotein c1/c2 (human)		D28382	77
ares parathyroid tumour (EST) (similar to human serum albumin pre	cursor)	AA860651	76
atagene mouse kidney (EST)	•	AA107327	75
ares parathyroid tumour NbHPA human cDNA (EST)	* *.	AA860653	74
ares mouse mammary gland (EST)	• • • •	AA619297	74
DNA clone 15 004 (EST) (human)		H01826	74
ares senescent fibroblasts (EST) (mouse)		W52190	74
eproalbumin (human)		E04315	72
DNA clone 73 169 (EST) (human)		T56624	72
amin D-binding protein (human)		L10641	71
oH gene (exon 8) (human)		Y11498	71
RL flow sorted chromosome	•	B05457	71
ares foetal liver spleen (EST) (mouse)		AA009524	71
ares foetal heart NbMH19W (EST) (mouse)		AA009421	69
ares foetal heart NbHH19W H. sapiens cDNA clone (EST)		W94377	67
enylalanine hydroxylase (human)		U49897	67
oline-5-carboxylate dehydrogenase (human)		U24266	66
utathione-S-transferase homologue (human)		U90313	65
I_CGAP_GCBI (EST) (human)		AA769294	65
otective protein (human)	•	M22960	64
one 27 375 (EST) (human)		N37046	62
atagene colon (#937 204) H. sapiens cDNA clone (EST)		AA149777	62
			

Refers to the nucleotide sequence homology between the cloned band isolated from the differential display and the correspondgene derived from the EMBL gene sequence bank.

Table 5

Identification of genes that were downregulated in male guinea / pig liver following 3-day treatment with WY-14,643

CASIM-EMBL gene	Accession No.	Sequences (%)
pig unless otherwise	าร เมณีโรกมาณและเลื่อง	the participation programming
stated) Ships to a definal	e definate regio	to proceed to the 1985
Complement C3 !	M34054	97 (5)
protein and protein	land Apple 51 Teach	September 18 Bright
Murinoglobulin	D84339	(95. 5 g)
Alpha-1-an-litter	M57274.	,
tiproteinase F	9740 10 (1111) - 3263345	egyani (ilgo rom)
Elongation lactor-ai-	A02243	89; · ; _ ; _ ;;
Counting protein G	V 04400	A Property of the Section of the Company
Coupinis protein Cr .	AUTTUZ	00
(human) NCI_CGAP_Ov1	A A 586300	07
(ECTP):(humon)	1 WW2000031 -11-	
Lecithin:cholesterol	D13668	1
acetyl transferase	 -	
(rabbit)	1.77	300
(rabbit) : Aldolase B (human)	X00270	ν.
Anti-thrombin III	E00116	80
	200110	
(human) Phenylalanine hy-	K03020	80
droxylase (human)	.1	* .
droxylase (human) Inter-α-trypsin in-	D38595	79
hibitor (human)	A contract of the contract of	
Normalised rat mus-	AA849753	78
:.cle (FST) (SS)		
Normalised rat ovary	AA801059	78
(FST) (ASS)		
Complement factor	X00284	77
Ba fragment (hu- man)	•	
man)		
Dihydrodiol dehydro-	U05598	76
PEHASE (IIIIIIIIIII)		•
Spot14 gene (thyroid-	Y08409	.75
inducible nepatic		· · · · · · · · · · · · · · · · · · ·
Pro 11111/(1111111111)		
BAC clone 174p12	AC004236	75
(human)	3/05/00	54
Mitochondrial alde-	X05409	74
hyde dehydroge-		
nase (human)	E04315	74
Preproalbumin (hu- man)	E04313	74
•	AA533142	74
NCI_CGAP_Pr9 (EST) (human) (S)	MM333142	74
Normalised rat pla-	AA851197	74
	ונוונטתח	'
centa (EXIII/AXI		
centa (EST) (AS)	104621	73
Heparin sulfate pro-	J04621	73
	J04621 R24330	73 73

Table 5 (Continued)

FASTA-EMBL gene identification (guineapig unless otherwise stated)		homolome (9/)
	والمراجع المعارب المعارب	<i>6</i> '-
Retinol dehydrogenase	. 033301	/1
(rat)		
TAPA-1 integral mem-	S45012	71
brane, protein	ri dirimahang meg	្សាទី"បាលខេត្ត។ សេចក្រ 👵
(CD81) (mouse) Complement component c5s	M35525	70
nent-c5s, 24	Transfer in the second	of Missional History (Park)
Apolinoprotein B. (nie)	5 - / J 11235	40
Apolipoprotein B (pig) cDNA clone 143 918 (EST) (human)	R76742	126 com Sale North
(EST) (human) α-fibrinogen (human) Soares foetal liver	ใช้ใช้ใช้ใช้เรื่องสำหรับก็ละ	THE SOURCE SHOW
α-fibringen (human)	K02560	20 Sinal hegieter
Soares foetal liver	W03726	20 Traction 99.
spleen INF (mouse)		00
spleen lNF (mouse) Barstead bowel (EST) (mouse)	A A 232040	67 F Mar R.J.
(mouse)	The Market of the	P 可量为14.6亿元。
(mouse) UDP glucuronosyl ransferase (cat)	A F0300137	tigal a movies made and
transferase (cat)	0303137.	20 CN (31-
Myeloid leukaemia cell	1.08246	65
Myeloid leukaemia cell. differentiation	3 - K	Straig Oaker Jan
differentiation protein (MCL-1)	totta karministi.	Commercial Commercial
protein (MCL-1) (human) (S)	'. 'n a 1	Fig. 1 1 - 39 401
(human) (S) STS SHGC-34 987 (human) (AS)	G27984	65 11 month of A.
man) (AS)		Superior Spires by
Soares mouse	AA222798	64
3NME125	the Landing of	
Stratagene mouse em-	AA199420	64************
bryonic (EST) (S)	i giri i <mark>Ti</mark> br çir sı	1 The state of the same
Rad 52 (mouse)	AF004854	63 : 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1

Refers to the nucleotide sequence homology between the cloned band isolated from the differential display and the corresponding gene derived from the EMBL gene sequence bank.

for several reasons: (1) the kinetics of ligation and transformation favour the isolation of smaller PCR products, thereby producing a misrepresentation of larger gene products; (2) northern blot analysis is notoriously insensitive and is unlikely to confirm expression of rare transcripts; (3) there is no measurable end point to the screening of clones produced in this way other than to analyse every transformed colony. We used instead an alternative approach; after running out the differ-

b EST is 'expressed sequence tag' — a gene of as yet unknown identity and function

Where sequence homologies were equal in both directions, boththe sense (S) and antisense (A) identities are given.

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ential display on a high-resolution agarose gel (Fig. 1) and overstaining with SYBR Green I to enhance visualisation, the composite bands were individually extracted, reamplified and cloned However, it has been well documented that single bands from differential displays often contain a heterogeneous mixture of different products (Mathieu-Daude et al., 1996; Smith et al., 1997). This is because polyacrylamide gels cannot discriminate between DNA sequences that differ in size by less than about 0.2% (Sambrook et al., 1989). High-resolution agarose gels such as those used in this work are even less sensitive, normally only discriminating products that differ in size by at least 1.5%. The use of the HA-red screening step enables resolution of identical or nearly idenical sequences based on their AT content (Wawer et al., 1995) and is sensitive down to < 1% difference. Furthermore, it is rapid, technically simple and does not require the use of radiolabels. Geisinger et al. (1997) originally demonstrated the isefulness of using HA-red to identify different products cloned from the same band of an RNA lifferential display experiment by simultaneously unning them in normal agarose (to discriminate y size) and in normal agarose containing HA-red to discriminate by AT content). We have found hat this approach is equally useful for identifying ifferent gene species cloned from the same band f our SSH display.

Diatchenko et al. (1996) reported that SSH is ighly efficient at producing differentially exressed gene species. However, we also included a econd screening step to further confirm that the ones isolated from the differential display were ideed differentially expressed. Duplicate dotblots f the candidate clones were blotted with the isplay from which they were originally isolated ad with the 'reverse subtraction' display. To ake the reverse-subtracted probe, the subtractive ybridisation step of the procedure was carried ut using the original tester cDNA as a driver, id the original driver cDNA as a tester. In this ay, clones that are false positives can be idenied through their presence in both blots. Such lse positives most commonly arise through havg a very high abundance in the initial sample or iusual hybridisation properties (Li et al., 1994).

radio de la Charlettal de Carregas Although the SSH method itself has been shown to be efficient, and despite the screening step that we included, there is an important caveat to bear in mind — namely that it is important that all clones be considered only as 'candidates' until the actual abundance of otheir mRNA is quantitated in treated and control samples. To-98 wards this end, we examined the expression of a limited number of clones using semi-quantitative, RT-PCR. Albumin was used as the reference gene as we have previously found that the expression of this gene does not appear to change with the treatment regime that we used (Fig. 4, and data not shown). There are a number of interesting points to note from our results. The first is the presence of genes that serve as appropriate positive controls in the upregulated and downregulated series. For example, in the rat it can be seen that CYP4AI expression increases 14-fold following treatment. Although CYP4AI mRNA expression levels following WY-14,643 treatment have not been previously reported in this model, the figure compares favourably with that recorded by Bell et al. (1991), who used RNAse-protection to quantitate CYP4A1 in rat liver following treatment with methylclofenapate, another PP. In addition, we also confirmed that the peroxisomal enoyl-CoA:hydratase-3-hydroxyacyl-CoA bifunctional enzyme is also upregulated 9-fold, in agreement with the findings of Chen and Crane (1992).

A number of genes were downregulated following Wy-14,643 exposure, including CYP2C11 expression. Corton et al. (1997) reported similar findings and suggested that this may in part explain why male rats exposed to Wy-14,643 and some other PPs have high serum estradiol levels, as estradiol is a substrate for CYP2C11. We have also shown that the expression of contrapsin-like protease inhibitor (CLPI) was downregulated by Wy-14,643. This has not previously been reported, and we suggest that it may be linked to a requirement for increased availability of amino acids to accommodate the hepatomegaly induced by treatment. Although little is known of the function of parathymosin-a, (zinc2+-binding protein) it has been shown to interact with the globular domain of histone H1, suggesting a role in histone function (Kondili et al., 1996). In contrast to the

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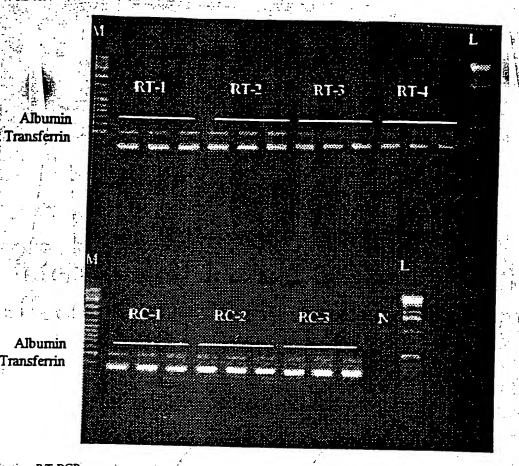


Fig. 4. Semi-quantitative RT-PCR experiment showing relative decrease in expression of transferrin in treated rat liver (RT-1 to RT-4) compared to controls (RC-1 to RC-3). An equal amount of mRNA was used in each reaction (10 ng), and each sample was quantitated in triplicate to reduce the effects of inter-tube variation. N is negative control (no mRNA). Lane M is a 100 bp ladder and lane L is a 1 Kb DNA ladder.

downregulation observed in this work, other studies have shown that parathymosin-a expression is elevated in breast cancer (Tsitsilonis et al., 1993, 1998), with the implication that parathymosin-α may somehow be involved in regulating cell proiferation by more than one mechanism. Transfer-∵in has previously been shown lownregulated in rat liver by hypolipidemic PPs Hertz et al., 1996). It is therefore interesting to note that we isolated a clone identified as transferin from the upregulated display profile. Since we confirmed by RT-PCR that transferrin is in fact lownregulated in the rat (Fig. 4), we conclude hat transferrin was either a false positive or was ncorrectly identified. It could also be that we ave isolated a close relative, splice variant or soform of transferrin, which demonstrates a diferent expression profile under these experimental onditions. Further investigations are therefore

required to determine which of these possibilities are correct.

One of our most intriguing observations was that one gene, CD81, appeared to be upregulated in rat liver but downregulated in guinea pig liver following Wy-14,643 exposure. CD81 is a widely expressed cell surface protein that is involved in a large number of cellular functions, including adhesion, activation, proliferation and differentiation (reviewed by Levy et al., 1998). Since all of these functions are altered to some extent in carcinogenesis, it is perhaps an important observation that CD81 expression is differentially regulated in a resistant and sensitive species exposed to a non-genotoxic carcinogen.

Albumin and ribosomal genes appear common to all differential displays and are thus undesirable false positives. However, due to their high expression in the liver, they are difficult to renove. We also noted a number of gene species, particularly in the guinea pig, which were common to both upregulated and downregulated profiles. Again, the most likely reason for these naving arisen is their high abundance.

A relatively large number of upregulated and lownregulated genes were isolated from guinea ig liver following Wy-14,643 exposure. However, he guinea pig genome has been relatively poorly haracterised and so many of the clones were dentified as resembling genes or ESTs from other pecies. Without full-length sequence data it is ifficult to ascertain the accuracy of the assigned ientities and this must be borne in mind when tilising data such as this, for example, in designng effective primers for RT-PCR studies. Alrough the actual isolated clone sequences can be sed to do this, their relatively small size often estricts the ability to design effective primers. In idition, as we observed with transferrin, using a ublished full-length sequence may help to idenfy false positives.

By comparing the expression profiles of genes showing altered expression in a PP-sensitive species (rat) with a PP-resistant species (guinea pig), it was our aim to identify genes that are mechanistically relevant to the non-genotoxic hepatocarcinogenic action of Wy-14,643. However, few of the genes that we have isolated were common to both the rat and the guinea pig. This suggests either that the molecular mechanisms of response in these two species are so different that few genes are commonly regulated in response to Wy-14,643 exposure, or that we have recovered only a small proportion of those genes that have altered expression. The latter seems the more likely scenario since it is perceived that one of the main problems of subtractive hybridisation and other differential expression technologies is the inability to consistently isolate rare gene transcripts (Bertioli et al., 1995). This is potentially problematic in that weakly expressed genes may play an important role in regulating key cellular processes, and that the majority of mRNA species are classified as

mi-quantitative RT-PCR analysis of selected gene species in the rat and guinea piga

Putative change o	f expression following ng to dotblot	Change according to RT-PCR quantitation	
Rat	Guinea pig	Rat	Guinea pig
N/A	N/A	No change	No al
Up	•		No change
Down	N/A	Downregulated*	N/O N/D
Up	N/A).//D
N/A	·		N/D
Up	Down	N/O	N/O Upregulated**(1.4
Down	N/A	Downregulated**	×) N/D
Down	N/A	Downregulated**	N/D
Up	N/A	Downregulated*	No change
Down	N/A	Downregulated**	N/O
Down	N/A		NID
Up			N/D N/O
	Rat N/A Up Down Up N/A Up Down Down Down Down Down Down Down	N/A N/A Up N/A Down N/A Up N/A N/A Up Up Down Down N/A Up N/A Down N/A Down N/A Down N/A Down N/A Down N/A	treatment according to dotblot Rat Guinea pig Rat N/A N/A No change Up N/A Upregulated* (9 ×) Down N/A Upregulated* (Abolished) Up N/A Up N/A Up No change Up N/A Up No change Up Down N/A Downregulated* (0.5 ×) Down N/A Downregulated** (0.6 ×) Down N/A Downregulated** (0.5 ×) Down N/A Downregulated** (0.5 ×) Down N/A Downregulated** (0.5 ×) Down N/A Downregulated* (0.5 ×) Down N/A Downregulated* (0.5 ×) N/A Downregulated* (0.5 ×) N/A Downregulated* (0.5 ×) Down N/A Downregulated* (0.5 ×) Down N/A Downregulated* (0.5 ×) N/A Downregulated*

N/A, not applicable; N/O, not optimised; N/D, not done. P < 0.0005;

^{*} P < 0.05.

trare in abundance (Bertioli et al., 1995). However, in their original paper describing the SSH technique, Gurskaya et al. (1996) demonstrated that SSH can enrich rare molecules between 1000and 5000-fold in a single round of hybridisation. Unfortunately, due to high background smearing in our initial experiments (which hindered identification of single bands), we were compelled to reduce the primary hybridisation time to only 4 h — a step that theoretically is likely to reduce the number of rare sequences (CLONTECHniques, CC 1996). Furthermore, it has been claimed by the manufacturers that, whilst this technique can identify changes as small as 1.5-fold between the driver and tester populations, it is best suited to the isolation of genes that show a greater than 5-fold increase (CLONTECHniques, 1996). In addition, where tester and driver contain genes with large and small differences in abundance, the SSH method will be biased towards identifying those genes with the large differences (CLONTECHniques, 1996). Thus, it is most probable that we have not isolated all of the more rarely expressed transcripts and those demonstrating small changes in expression.

One problem that remains is identifying the function of genes isolated in SSH experiments as described herein, some of which may be crucial to the process of carcinogenesis, and are, to date, unidentified. However, we have provided evidence herein that SSH can be used to begin the process of characterising the extent and importance of altered gene expression in response to a chemical stimulus. The developments of this approach should include characterisation of temporal and dose responses, and functional analysis studies including knockout mice. In combination, such studies should make a significant contribution to our understanding of the molecular mechanisms of action and physiological relevance of gene regulation in non-genotoxic hepatocarcinogenesis. It should then be possible to ascertain whether differentially expressed genes are causally or casually related to the chemical-induced toxicity, and therefore a substantial mechanistic advance.

It is clear that there are also broader applications for this experimental approach that go beyond understanding the molecular mechanisms of

peroxisome-proliferator induced non-genotoxic hepatocarcinogenesis in rodents. The potential medical and therapeutic benefits of elucidating the molecular changes that occur in any given cell in progressing from the normal to the carcinogenic (or other diseased, abnormal or developmental) state are very substantial. Notwithstanding the lack of complete functional identification of altered gene expression, such gene profiling studies described herein essentially provides a 'fingerprint' of each stage of carcinogenesis, and should help in the elucidation of specific and sensitive biomarkers for different types of cancer. Amongst other benefits, such fingerprints and biomarkers could help uncover differences in histologically identical cancers, and provide diagnostic tests for the earliest stages of neoplasia. In addition, the genes identified by this approach may be incorporated into gene-chip DNA-arrays, thus providing a standard genetic fingerprint for a particular toxin treatment in a particular species. Interrogation of these gene arrays for an unknown compound that has a similar pattern to the known reference chemical would then provide evidence that the unknown may have a toxicity profile similar to the 'standard' fingerprint, thereby serving as a mechanistically relevant platform for further detailed investigations.

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References

Anderson, N.L., Esquer-Blasco, R., Richardson, F., Foxworthy, P., Eacho, P., 1996. The effects of peroxisome proliferators on protein abundances in mouse liver. Toxicol. Appl. Pharmacol. 137, 75-89.

Ashby, J., 1992. Prediction of non-genotoxic carcinogenesis. Toxicol. Lett. 64-65, 605-612.

Bell, D.R., Bars, R.G., Gibson, G.G., Elcombe, C.R., 1991. Localisation and differential induction of cytochrome P450IVA and acyl coA oxidase in rat liver. Biochem. J. 275, 247-252.

Bertioli, D.J., Schlichter, U.H.A., Adams, M.J., Burrows, P.R., Steinbiss, H.-H., Antoniw, J.F., 1995. An analysis of

- number mRNAs. Nucleic Acid Res. 23 (21), 4520-4523.
 - Cattley, R.C., Kato, M., Popp, J.A., Teets, V.J., Voss, K.S., 1994. Initiator-specific promotion of hepatocarcinogenesis by Wy-14,643 and clofibrate. Carcinogenesis 15 (8), 1763-1766.
- Chen, N., Crane, D.I., 1992. Induction of the major integral in a membrane protein of mouse liver peroxisomes by peroxisome proliferators. Biochem J. 283, 605-610.
 - CLONTECHniques, 1996. Technical Tips: Clontech PCR-Select cDNA Subtraction, October 25, application notes.
 - CLONTECHniques, 1997. PCR-Select Differential Screening Kit The Next Step After Clontech PCR-Select cDNA Subtraction. XII(2), 18-19, application notes.
- Corton, J.C., Bocos, C., Moreno, E.S., Merrit, A., Cattley, R.C., Gustaffson, J.A., 1997. Peroxisome proliferators alter the expression of estrogen-metabolising enzymes. Biochimie 79, 151-162.
- Davis, M., Cohen, D.I., Nielson, E.A., Steinmetz, M., Paul, W.E., Hood, L., 1984. Cell-type-specific cDNA probes and the murine I region: the localisation and orientation of Ad/a. Proc. Natl. Acad. Sci USA 81, 2194-2198.
- Diatchenko, L., Lau, Y.-F.C., Campbell, A.P., Chenchik, A., Moqadam, F., Huang, B., Lukyanov, K., Gurskaya, N., Sverdlov, E.D., Siebert, P.D., 1996. Suppression subtractive hybridisation: a method for generating differentially regulated or tissue-specific cDNA probes and libraries. Proc. Natl. Acad. Sci. USA 93, 6025-6030.
- Duguid, J., Dinauer, M., 1990. Library subtraction of in vitro cDNA libraries to identify differentially expressed genes in scrapie infection. Nucleic Acid Res. 18 (9), 2789-2792.
- Geisinger, A., Rodriguez, R., Romero, V., Wettstein, R., 1997.

 A simple method for screening cDNAs arising from the cloning of RNA differential display bands. Elsevier trends journals technical tips online, http://tto.trends.com, document number T01110
- Guimaraes, M.J., Lee, F., Zlotnik, A., McClanahan, T., 1995.
 Differential display by PCR: novel findings and applications. Nucleic Acid Res. 23 (10), 1832-1833.
- Gurskaya, N.G., Diatchenko, L., Chenchik, P.D., Siebert, P.D., Khaspekov, G.L., Lukyanov, K.A., Vagner, L.L., Ermolaeva, O.D., Lukyanov, S.A., Sverdlov, E.D., 1996. Equalising cDNA subtraction based on selective suppression of polymerase chain reaction: cloning of Jurkat cell transcripts induced by phytohemaglutinin and phorbol 12-myrystate 13-acetate. Anal. Biochem. 240, 90-97.
- Hayashi, F., Tamura, H., Yamada, J., Kasai, H., Suga, T., 1994. Characteristics of the hepatocarcinogenesis caused by dehydroepiandrosterone, a peroxisome proliferator, in male F-344 rats. Carcinogenesis 15 (190), 2215-2219.
- Hedrick, S.M., Cohen, D.I., Nielsen, E.A., Davis, M.M., 1984.
 Isolation of cDNA clones encoding T cell-specific membrane-associated proteins. Nature 308 (8), 149-153.
- Hertz, R., Seckbach, M., Zakin, M.M., Bar-Tana, J., 1996. Transcriptional suppression of the transferin gene by hypolipidemic peroxisome proliferators. J. Biol. Chem. 271 (1), 218-224.

- Hubank, M., Schatz, D.G., 1994. Identifying differences in mRNA expression by representational difference analysis. Nucleic Acid Res. 22 (25), 5640-5648.
- Human and Experimental Toxicology, 1994, Hum. Exp. Toxicol. 13 (Suppl. 2) (entire issue).
- Kolls, J., Dsininger, P., Cohen, C., Larson, J., 1993. cDNA equalisation for reverse transcription-polymerase chain reaction quantitation. Anal. Biochem 208, 264-269.
- Kondili, K., Tsolas, O., Papamarcaki, T., 1996, Selective interaction between parathymosin and histone H1. Eur. J. Biochem. 242 (1), 67-74.
- Lake, B.G., Evans, J.G., Gray, T.J.B., Korosi, S.A., North, C.J., 1989. Comparative studies of nafenopin-induced hepatic peroxisome proliferation in the rat, Syrian hamster, guiea pig and marmoset. Toxicol. Appl. Pharmacol. 99, 148-160.
- Lake, B.G., Evans, J.G., Cunninghame, M.E., Price, R.J., 1993. Comparison of the hepatic effects of Wy-14,643 on peroxisome proliferation and cell replication in the rat and Syrian hamster. Environ. Health Perspect. 101 (S5), 241-248.
- Levy, S., Todd, S.C., Maecker, H.T., 1998. CD81 (TAPA-1): a molecule involved in signal transduction and cell adhesion in the immune system. Annu. Rev. Immunol. 16, 89-109.
- Li, W.B., Gruber, C.E., Lin, J.J., D'Alessio, J.M., Jessee, J.A., 1994. The isolation of differentially expressed genes in fibroblast growth factor stimulated BC3H1 cells by subtractive hybridization. BioTechniques 16, 722-729.
- Liang, P., Pardee, A.B., 1992. Differential display of eukaryotic messenger RNA by means of the polymerase chain reaction. Science 257 (5072), 967-971.
- Liang, P., Averboukh, L., Keyomarsi, K., Sager, R., Pardee, A.B., 1992. Differential display and cloning of messenger RNAs from human breast cancer versus mammary epithelium. Cancer Res. 52, 6966-6968.
- Lipman, D.J., Pearson, W.R., 1985. Rapid and sensitive protein similarity searches. Science 227, 1435-1441.
- Makowska, J.M., Gibson, G.G., Bonner, F.W., 1992. Species differences in ciprofibrate induction of hepaic cytochrome P450IVA1 and peroxisome proliferation. J. Biochem. Toxicol. 7, 183-191.
- Marsman, D.S., Cattley, R.C., Conway, J.G., Popp, J.A., 1988. Relationship of hepatic peroxisome proliferation and replicative DNA synthesis to the hepatocarcinogenicity of the peroxisome proliferators di-(2-ethylhexyl)phthalate and [4-chloro-6-(2,3-xylidino)-2-pyrimidinylthio]acetic acid (Wy-14,643) in rats. Cancer Res. 48, 6739-6744;
- Mathieu-Daude, F., Cheng, R., Welsh, J., McClelland, M., 1996. Screening of differentially amplified cDNA products from RNA arbitrarily primed PCR fingerprints using single strand conformation polymorphism (SSCP) gels. Nucleic Acid Res. 24 (8), 1504-1507.
- Orton, T.C., Adam, H.K., Bentley, M., Holloway, B., Tucker, M.J., 1984. Clobuzarit: species differences in the morphological and biochemical response of the liver following chronic administration. Toxicol. Appl. Pharmacol. 73, 138-151.

- Parodi, S., 1992. Non-genotoxic factors in the carcinogenic property process: problems of detection and hazard evaluation.

 WAS ATOXICOL Lett. 64-65, 621-630, Pariotoxical
- Pearson, W.R., Lipman, D.J., 1988. Imported tools for biological sequence comparison. Proc. Natl. Acad. Sci. USA 85, 2444-2448.
- Rockett, J.C., Esdaile, D.J., Gibson, G.G., 1997. Molecular profiling of non-genotoxic hepatocarcinogenesis using differential display reverse transcription-polymerase chain reaction (ddRT-PCR). Eur. J. Drug. Metab. Pharmacokinet 22.(4), 329-333.
- Rodricks, J.V., Turnbull, D., 1987. Inter-species differences in peroxisomes and peroxisome proliferation. Toxicol: Ind. Health 3, 197-212.
- Sambrook, J., Fritsch, E.F., Maniatis, T., 1989. In: Ford, N., Nolan, C., Ferguson, M. (Eds.), Molecular Cloning A Laboratory Manual, second ed. Cold Spring Harbor Laboratory Press, New York.
 - Sargent, T., Dawid, I., 1983. Differential gene expression in the gastrula of *Xenopus laevis*. Science 222, 135-139.
 - Smith, N.R., Li, A., Aldersley, M., High, A.s., Markham, A.F., Robinson, P.A., 1997. Rapid determination of the

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- complexity of cDNA bands extracted from DDRT-PCR polyacrylamide gels. Nucleic Acid Res. 25 (17), 3552-3554.
- Tsitsilonis, O.E., Stiakakis, J., Koutselinis, A., Gogas, J., Markopoulos, C., Yialouris, P., Bekris, S., Panoussopoulos, D., Kiortsis, V., Voelter, W., Haritos, A.A., 1993.

 Expression of alpha-thymosins in human tissues in normal and abnormal growth. Proc. Natl. Acad. Sci. USA 90 (20), 9504-9507.
- Tsitsilonis, O.E., Bekris, E., Voutsas, I.F., Baxevanis, C.N., Markopoulos, C., Papadopoulou, S.A., Kontzoglou, K., Stoeva, S., Gogas, J., Voelter, W., Papamichail, M., 1998.

 The prognostic value of alpha-thymosins in breast cancer.

 Anticancer, Res. 18 (3A), 1501-1508.
- Wan, J.S., Sharp, S.J., Poirier, G.M.-C., Wagaman, P.C., Chambers, J., Pyati, J., Hom, Y.-L., Galindo, J.E., Huvar, A., Peterson, P.A., Jackson, M.R., Erlander, M.G. 1996. Cloning differentially expressed mRNAs. Nat. Biotechnol. 14, 1685-1691.
- Wawer, C.; Ruggeberg, H., Meyer, G., Muyzer, G., 1995. A simple and rapid electrophoresis method to detect sequence variation in PCR-amplified DNA fragments. Nucleic Acid Res. 23 (23), 4928-4929.

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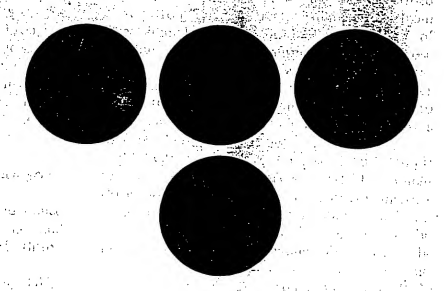
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Yeast microarrays for genome wide parallel genetic and gene of the first of promoderny pathogs. expression analysis

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Contributed by Ronald W. Davis, September 2, 1997

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ABSTRACT We have developed high-density DNA microarrays of yeast ORFs. These microarrays can monitor hybridization to ORFs for applications such as quantitative differential gene expression analysis and screening for sequence polymorphisms. Automated scripts retrieved sequence information from public databases to locate predicted ORFs and select appropriate primers for amplification. The primers were used to amplify yeast ORFs in 96-well plates, and the resulting products were arrayed using an automated micro arraying device. Arrays containing up to 2,479 yeast ORFs were printed on a single slide. The hybridization of fluorescently labeled samples to the array were detected and quantitated with a laser confocal scanning microscope. Applications of the microarrays are shown for genetic and gene expression analysis at the whole genome level.

The genome sequencing projects have generated and will continue to generate enormous amounts of sequence data. The genomes of Saccharomyces cerevisiae, Haemophilus influenzae (1), Mycoplasma genitalium (2), and Methanococcus jannischii (3) have been completely sequenced. Other model organisms have had substantial portions of their genomes sequenced as well including the nematode Caenorhabditis elegans (4) and the small flowering plant Arabidopsis thaliana (5). Given this everincreasing amount of sequence information, new strategies are necessary to efficiently pursue the next phase of the genome projects—the elucidation of gene expression patterns and gene product function on a whole genome scale.

One important use of genome sequence data is to attempt to identify the functions of predicted ORFs within the genome. Many of the ORFs identified in the yeast genome sequence were not identified in decades of genetic studies and have no significant homology to previously identified sequences in the database. In addition, even in cases where ORFs have significant homology to sequences in the database, or have known sequence motifs (e.g., protein kinase), this is not sufficient to determine the actual biological role of the gene product. Experimental analysis must be performed to thoroughly understand the biological function of a given ORF's product. Model organisms, such as S. cerevisiae, will be extremely important in improving our understanding of other more complex and less manipulable organisms.

To examine in detail the functional role of individual ORFs and relationships between genes at the expression level, this work describes the use of genome sequence information to study large numbers of genes efficiently and systematically. The procedure was as follows. (i) Software scripts scanned annotated sequence information from public databases for predicted ORFs. (ii) The start and stop position of each identified ORF was extracted automatically, along with the sequence data of the ORF and 200

bases flanking either side. (iii) These data were used to automatically select PCR primers that would amplify the ORF. (iv) The primer sequences were automatically input into the automated multiplex oligonucleotide synthesizer (6). (v) The oligonucleotides were synthesized in 96-well format, and (vi) used in 96-well format to amplify the desired ORFs from a genomic DNA template. (vii) The products were arrayed using a high-density DNA arrayer (7-10). The gene arrays can be used for hybridization with a variety of labeled products such as cDNA for gene expression analysis or genomic DNA for strain comparisons, and genomic mismatch scanning purified DNA for genotyping (11).

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METHODS

Script Design. All scripts were written in UNIX Tool Command Language. Annotated sequence information from GenBank was extracted into one file containing the complete nucleotide sequence of a single chromosome. A second file contained the assigned ORF name followed by the start and stop positions of that ORF. The actual sequence contained within the specified range, along with 200 bases of sequence flanking both sides, was extracted and input into the primer selection program PRIMER 0.5 (Whitehead Institute, Boston). Primers were designed so as to allow amplification of entire ORFs. The selected primer sequences were read by the 96-well automated multiplex oligonucleotide synthesizer instrument for primer synthesis. The forward and reverse primers were synthesized in two separate 96-well plates in corresponding wells. All primers were synthesized on a 20-nmol scale.

ORF Amplification and Purification. Genomic DNA was isolated as described (12) and used as template for the amplification reactions. Each PCR was done in a total volume of 100 µl. A total of 0.2 μ M each of forward and reverse primers were aliquoted into a 96-well PCR plate (Robbins Scientific, Sunnyvale, CA); a master mix containing 0.24 mM each dNTP, 10 mM Tris (pH 8.5), 50 mM MgCl₂, 2.5 units Taq polymerase, and 10 ng of template was added to the primers, and the entire mix was thermal cycled for 30 cycles as follows: 15 min at 94°C, 15 min at 54°C, and 30 min at 72°C. Products were ethanol precipitated in polystyrene v-bottom 96well plates (Costar). All samples were dried and stored at -20°C.

Arraying Procedure and Processing. Microarrays were made as described (8).

A custom built arraying robot was used to print batches of 48 slides. The robot utilizes four printing tips which simultaneously pick up ~1 µl of solution from 96-well microtiter plates. After printing, the microarrays were rehydrated for 30 sec in a humid chamber and then snap dried for 2 sec on a hot plate (100°C). The DNA was then UV crosslinked to the surface by subjecting the slides to 60 millijoules of energy. The rest of the poly-L-lysine surface was blocked by a 15-min incubation in a solution of 70 mM succinic anhydride dissolved in a solution consisting of 315 ml of 1-methyl-2-pyrrolidinone (Aldrich) and 35 ml of 1 M boric acid (pH 8.0). Directly after the blocking reaction, the bound DNA was denatured by a 2-min incubation in distilled water at ~95°C.

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Abbreviation: YEP, yeast extract/peptone.

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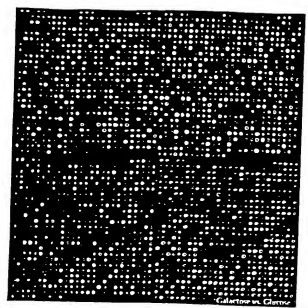


Fig. 1. Two-color fluorescent scan of a yeast microarray containing 2,479 elements (ORFs). The center-to-center distance between elements is 345 μ m. A probe mixture consisting of cDNA from yeast extract/peptone (YEP) galactose (green pseudocolor) and YEP glucose (red pseudocolor) grown yeast cultures was hybridized to the array. Intensity per element corresponds to ORF expression, and pseudocolor per element corresponds to relative ORF expression between the two cultures.

The slides were then transferred into a bath of 100% ethanol at room temperature.

Probe Preparation: cDNA. Yeast cultures (100 ml) were grown to ~1 OD_{A600} and total RNA was isolated as described (13). Up to 500 μg total RNA was used to isolate mRNA (Qiagen, Chatsworth, CA). Oligo(dT)20 (5 μg) was added and annealed to 2 μg of mRNA by heating the reaction to 70°C for 10 min and quick chilling on ice, plus 2 μl SuperScript II (200 units/μl) (Life Technologies, Gaithersburg, MD), 0.6 μl 50× dNTP mix (final concentrations were 500 μM dATP, dCTP, dGTP, and 200 μM dTTP), 6 μl 5× reaction buffer, and 60 μM Cy3-dUTP or Cy5-dUTP (Amersham). Reactions were carried out at 42°C for 2 h, after which the mRNA was degraded by the addition of 0.3 μl 5 M NaOH and 0.3 μl 100 mM EDTA and heating to 65°C for 10 min. The sample was then diluted to 500 μl with TE and concentrated using a Microcon-30 (Amicon) to 10 μl.

Probe Preparation: Genomic DNA. Fluorescent DNA was prepared from total genomic DNA as follows: 1 µg of random nonamer oligonucleotides was added to 2.5 µg of genomic DNA. This mixture was boiled for 2 min and then chilled on ice. A reaction mixture containing dNTPs (25 µM dATP, dCTP, dGTP, 10 µM dTTP, and 40 µM Cy3-dUTP or Cy5-dUTP) reaction buffer (New England Biolabs), and 20 units exonuclease free Klenow enzyme (United States Biochemical) was added, and the reaction was incubated at 37°C for 2 h. The sample was then diluted to 500 µl with TE and concentrated using a Microcon-30 (Amicon) to 10 µl.

Hybridization. Purified, labeled probe was resuspended in 11 μ l of 3.5× SSC containing 10 μ g Escherichia coli tRNA, and 0.3% SDS. The sample was then heated for 2 min in boiling water, cooled rapidly to room temperature, and applied to the array. The array was placed in a sealed, humidified, hybridization chamber. Hybridization was carried out for 10 h in a 62°C water bath, after which the arrays were washed immediately in 2× SSC/0.2% SDS. A second wash was performed in 0.1× SSC.

Analysis and Quantitation. Arrays were scanned on a scanning laser fluorescence microscope developed by Steve Smith with software written by Noam Ziv (Stanford Univer-

sity). A separate scan was done for each of the two fluorophores used. The images were then combined for analysis. A bounding box, fitted to the size of the DNA spots, was placed over each array element. The average fluorescent intensity was calculated by summing the intensities of each pixel present in a bounding box and then dividing by the total number of pixels. Local area background was calculated for each array element by determining the average fluorescent intensity at the edge of the bounding box. To normalize for fluorophore-specific variation, control spots containing yeast genomic DNA were applied to each quadrant during the arraying process. These elements were quantitated and the ratios of the signals were determined. These ratios were then used to normalize the photomultiplier sensitivity settings such that the ratios of the fluorescence of the genomic DNA spots were close to a value of 1.0. The average signal intensity at any given spot was regarded as significant if it was at least two standard deviations above background. Each experiment was conducted in duplicate, with the fluorophores representing each channel reversed. The ratios presented here are the average of the two experiments, except in the case in which the signal for the element in question was below the reliability threshold. The reliability threshold also determined the dynamic range of the experiment. For all of the experiments presented, the average dynamic range was ~1 to 100. In the case where the fluorescence from a very bright spot saturates the detector, differential ratios will, in general, be underestimated. This can be compensated for by scanning at a lower overall sensitivity.

RESULTS

The accumulation of sequence information from model organisms presents an enormous opportunity and challenge to understand the biological function of many previously uncharacterized genes. To do this accurately and efficiently, a directed strategy was developed that enables the monitoring of multiple genes simultaneously. Microarraying technology provides a method by which DNA can be attached to a glass surface in a high-density format (8). In practice, it is possible to array over 6,000 elements in an area less than 1.8 cm². Given that the yeast genome consists of ~6,100 ORFs, the entire set of yeast genes can be spotted onto a single glass slide.

With this capability and the availability of the entire sequence of the yeast genome, our strategy was to use a directed approach for generating the complete genome array. This procedure involved synthesizing a pair of oligonucleotide primers to amplify each ORF. The PCR product containing each gene of interest was arrayed onto glass and used, for example, as probe for monitoring gene expression levels by hybridizing to the array labeled cDNA generated from isolated mRNA of a culture grown under any experimental condition.

Primer Selection and Synthesis. The primer selection was fully automated using Tool Command Language scripts and PRIMER 0.5. (Whitehead). Primer pairs were automatically selected successfully for >99% of the ORFs tested. Primer sequences can thus be selected rapidly with minimal manual processing. A complete set of forward and reverse primers were selected initially for each ORF on chromosomes I, II, III, V, VI, VIII, IX, X, and XI. Primers for a representative set of ORFs (15% coverage) were chosen for the remaining chromosomes. With the release of the entire yeast genome sequence, the complete set of primers has now been selected.

Because each ORF requires a unique pair of synthetic primers, a total of approximately 12,200 oligonucleotides will be required to individually amplify each target. This costly component was addressed with the automated multiplex oligonucleotide synthesizer (6) which efficiently synthesizes primers in a 96-well format. Each primer, synthesized on a 20-nmol scale, provides enough material for 100 amplification reactions, whereas a given PCR product provides enough material to generate an element on

Table 1. Heat shock vs. control expression data

Ratio gene expr			, t	and the state of t
Control	Heat	ORF	Gene	Description
4	2.2	YLR142	PUT1	Proline oxidase
,	2.0	YOL140	ARG8	Acetylomithine aminotransferase
23	•	YGL148:	. ARO2	Chorismate synthase
	36.0	YFL014	HSP12	Heat shock protein
;	27.4	YBR072	HSP26	Heat shock protein
	6.7	YBR054	YRO2	Similarina a Troppe a
	3.4	YCR021	HSP30	Similarity to HSP30 heat shock protein Yrolp
	2.6	YER103	SSA4	Heat shock protein
11.70	.25	YLR259	HSP60	Heat shock protein
	2.1.	YBR169	SSE2	Mitochondrial heat shock protein HSP60
·	1.7	YBL075	SSA3	Heat shock protein of the HSP70 family
	1.4	YPL240	HSP82	Cytoplasmic heat shock protein
	1.4	YDR258	HSP78	Heat shock protein
1.0	٠٠	YNL007	SISI	Mitochondrial heat shock protein of clpb family of ATP-dependent protes Heat shock protein
1.1		YEL030	0101	
1.9		YHR064		70-kDa heat shock protein
	1.3	YBL008	HIR1	Heat shock protein
2.6		YBL002	HTB2	Histone transcription regulator
3.3	3	YBL003	HTA2	Histone H2B.2
3.3	**	YBR010	HHT1	Histone H2A.2
3.9 ·		YBR009	HHF1	Histone H3
	- 2.4	YDR343	НХТ6	Histone H4
	2:1	YHR092	HXT4	High-affinity hexose transporter
.6		YAR071 .	PHO11	Moderate- to low-affinity glucose transporter
. 7.	2.3	YLR096	KIN2	Secreted acid phosphatase, 56 kDa isozyme
. 5		YER102 -	RPS8B	Ser/ In protein kinase
.6	•	YBR181	RPS101	Ribosomal protein S8.e
.6		YCR031	CRY1	Ribosomal protein S6.e
.7		YLR441	RP10A	40S ribosomal protein S14.e
.8	•	YHR141	RPL41B	Ribosomal protein S3.a.e
.8		YBL072	RPS8A	Ribosomal protein L36a.e
.8		YHL015	URP2	Ribosomal protein S8.e
8	•	YBR191	URP1	Ribosomal protein
.1		YLR340	RPLA0	Ribosomal protein L21.e
3 .		YGL123	SUP44	Acidic Ribosomal protein L10.e
	5.8	YLR194	-3. **	Ribosomal protein
				Hypothetical protein

500-1,000 arrays. Thus, a single primer pair provides enough starting material for up to ~50,000 arrays.

Primers were synthesized to amplify yeast ORFs. Primer synthesis had a failure rate of <1% in over 18 plates of synthesis as determined by standard trityl analysis (6). The success rate of the PCR amplifications using the primer pairs was 94% based on agarose gel analysis of each PCR. The purified PCR products were used to generate arrays. Two versions of the arrays were created for the experimental results presented here. The first array contained 2,287 elements and the second array batch contained 2,479 elements.

Genome Arrays. The amplified ORFs were arrayed onto glass at a spacing of 345 microns (Fig. 1). The high-density spacing of DNA samples allows the hybridization volumes to be minimized—volumes are a maximum of $10~\mu$ l. The labeled probe can thus be maintained at relatively high concentrations, making $1-2~\mu$ g of mRNA sufficient for analysis. This also obviates the need for a subsequent amplification step and thus avoids the risk of altering the relative ratios of different cDNA species in the sample.

Genetic Analysis: Genomic Comparison of Unrelated Strains. Microarrays allow efficient comparison of the genomes of different strains. Genomic DNA from Y55, an S. cerevisiae strain divergent from the reference strain S288c, was randomly labeled with Cy3-dUTP and hybridized simultaneously with the S288c DNA labeled with Cy5-dUTP. When a comparison between the hybridization of the DNA from the two strains was done, several

elements gave relatively little or no signal above background from the Cy3 channel (data not shown). These include SGE1, ASP3A-D, YLR156, YLR159, YLR161, ENA2 (YDR039 is ENA2), and YCR105. These results imply that the regions containing these genes are extremely divergent, or all together deleted from the strain. Subsequent attempts to generate PCR products from SGE1, ENA2, and ASP3A using Y55 DNA failed. This result supports the conclusion that these genes are likely to be missing from the Y55 genome. It is interesting to note that at least two of the regions absent in the Y55 genome have been previously shown or suggested to be deleted in mutant laboratory strains (14-16). In particular, the Asp-3 region appears to be highly prone to being deleted (15, 16).

These results indicate that gene arrays can be used to efficiently screen different strains of an organism for large deletion polymorphisms. A single hybridization and scan will reveal differences based on differential hybridization to particular elements. It is reasonable to suppose that an equivalent number of genes are present in the Y55 genome and absent in the S288c genome. This result should be viewed as a minimum estimate of the deletion polymorphisms that exist between these two unrelated strains as intergenic deletions or small intragenic deletions would not be detected because considerable hybridizing material would be remain. Sequence polymorphisms, such as deletions, are present in populations of every species and must at some level affect phenotype. One of the challenges of the genome era will be to critically examine sequence polymorphisms that exist in the natural gene pool relative to the reference genome sequence.

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Fig. 2. ORF categories displaying differential expression between heat shocked and untreated cultures. Bars within categories correspond to individual ORFs. Green shaded bars correspond to relative increases in ORF expression under 25°C growth conditions. Red shaded bars correspond to relative increases in ORF expression under 39°C growth conditions.

Gene Expression Analysis. The arrays were used to examine gene expression in yeast grown under a variety of different conditions. Expression analysis is an ideal application of these arrays because a single hybridization provides quantitative expres-

sion data for thousands of genes. To better understand results for genes of known function, ORFs were placed in biologically relevant categories on the basis of function (e.g., amino acid catabolic genes) and/or pathways (e.g., the histidine biosynthesis pathway).

Table 2. Cold shock vs. control expression data

Ratio of gene expression		1	
Control Cold	ORF	Gene	Description
2.4 2.9 1.4 2.0 2.1 2.2 2.7 3.1 3.3 3.8 7.9 7.9 16.5 1.8 2.1 2.4 2.8 3.4 2.3 1.7 1.7 1.6 1.6 3.1 2.2 2.0 4.2	YOR153 YCR012 YCL040 YHR064 YJL034 YDR258 YLL039 YLL026 YER103 YBR126 YPL240 YBR054 YBR072 YCR021 YDR343 YHR096 YFR053 YHR096 YFR053 YHR092 YHR094 YHR089 YLR048 YLR441 YLL045 YLR029 YGL123 YBR067 YER011 YCR058 YKL102	PDR5 PGK1 GLK1 KAR2 HSP78 UB14 HSP104 SSA4 TPS1 HSP82 YRO2 HSP26 HSP26 HSP30 HXT6 HXT5 HXK1 HXT4 HXT1 GAR1 NAB1B RP10A RPL4B RPL13A SUP44 TIP1 TIR1	Pleiotropic drug resistance protein Phosphoglycerate kinase Aldohexose specific glucokinase Heat shock protein Nuclear fusion protein Mitochondrial heat shock protein of clpb family of ATP-dependent proteases Ubiquitin precursor Heat shock protein Heat shock protein Heat shock protein Similarity to HSP30 heat shock protein Yrolp Heat shock protein House shock protein Heat shock protein House finity hexose transporter Putative hexose transporter Hexokinase I Moderate- to low-affinity glucose transporter Low-affinity hexose (glucose) transporter Nucleolar rRNA processing protein 40S ribosomal protein p40 homolog b Ribosomal protein S3a.c Ribosomal protein L15.e Ribosomal protein L15.e Ribosomal protein L15.e Ribosomal protein Cold- and heat-shock-induced protein of the Srp1/Tip1p family Cold-shock-induced protein of the Tir1p, Tip1p family

Table 3. Glucose vs. galactose expression data 4.1

	io of pression			
Glucose	Galactose	ORF ORF	Gene	George Description (p. 6)
2.1	2.	YHR018	ARG4	Arginosuccinate lyase
3.5	2.5	YPR035	GLN1	Glutamate-ámmonia ligase (1994) (1995)
2.8		YML116	ATR1	Aminotriazole and 4-nitroquinoline resistance protein
2.0	ί.	YMR303	ADH2	Alcohol dehydrogenase II
3.7		YBR145	ADH5	Alcohol dehydrogenase V
	3.2	YBL030	AAC2	ADP, ATP carrier protein 2 in additional products Viole
	2.9	YBR085	AAC3	ADP, ATP carrier protein
	2.7	YDR298	ATP5	H ⁺ -transporting ATP synthase δ chain precursor
•	25	YBR039 1250	ATP3	H+-transporting ATP synthase y chain precursor
	5.5	YML054	CYB2	Lactate dehydrogenase cytochrome b211.57 17.300
	3.4	YML054	CYB2	Lactate dehydrogenase cytochrome b2
	2.3	YKL150	MCR1	Cytochrome-65 reductase
	4.2	YBL045	COR1	I Thigning of greathern in and warm 4425 and a section in
	3.5	YDL067 "	_	Cytochrome e oxidase chain VIIA
; '	2.7	YLR038	COX12	Cytochrome c oxidase, subunit VIB
	2.6	YHR051	COX6	Cytochrome c oxidase subunit VI
,	2.4	YLR395	COX8	Cytochrome'e oxidase chain VIII
	2.3	YFR033	QCR6	Ubiquinol-cytochrome c reductase 17K protein
	23.7	YLR081	GAL2	Galactose (and glucose) permease
•	21.9	YBR018	GAL7	UDP-glucose-hexose-1-phosphate uridylyltransferase
	21.8	YBR020	GAL1	Galactokinase
	19.5	YBR019	GAL10	UDP-glucose 4-epimerase
	14.7	YLR081 ^	GAL2	Galactose (and glucose) permease
	8.6	YDR009	GAL3	Galactokinase
	3.0	YML051 (GAL80(1)	Negative regulator for expression of galactose-induced genes
	2.8		GAL80(2)	Negative regulator for expression of galactose-induced genes
2.7		YER055	HIS1	ATP phosphoribosyltransferase
3.4		YBR248	HIS7	Glutamine amidotransferase/cyclase
,			• •	Phosphoribosyl-AMP cyclohydrolasc/phosphoribosyl-ATP pyrophosphatase/histidinol
7.4		YCL030	HIS4	dehydrogenase dehydrogenase
5.8		YKR080	MTD1	Methylenetetrahydrofolate dehydrogenase (NAD+)
6.0		YDR019	GCV1	Glycine decarboxylase T subunit
6.1		YLR058 '	SHM2	Serine hydroxymethyltransferase
	8.1	YML123	PHO84	High-affinity inorganic phosphate/H+ symporter
3.5		YDR408	ADE8	Phosphoribosylglycinamide formyltransferase (GART)
3.6		YDR408	ADE8	Phosphoribosylglycinamide formyltransferase (GART)
4.4		YAR015	ADE1	Phosphoribosylamidoimidazole-succinocarboxamide synthase
5.6		YMR300	ADE4	Amidophosphoribosyltransferase
5.6		YOR128	ADE2	Phosphoribosylaminoimidazole carboxylase
6.0:			ADE5,7	Phosphoribosylamine-glycine ligase and phosphoribosylformylglycinamidine cyclo-ligase
	6.3	YBL015	ACH1	Acetyl-CoA hydrolase

Heat Shock Results. A log phase culture growing in YEP/ dextrose medium at 25°C was split in half. One half of the culture remained at 25°C whereas the other half of the culture was shifted to 39°C. mRNA was isolated from both cultures 1 h after heat shock for comparison on microarrays and, although this time point is not optimal for measuring induction of heat shock mRNAs (17), many known heat shock genes exhibited considerable induction at this time point (Table 1; Fig. 2). Down-regulation of genes in the ribosomal protein and histone gene categories was also observed. Differential expression between the heat-shocked culture and the control was also observed for many other genes. Genes in many categories, such as amino acid catabolism and amino acid synthesis, exhibited a mixed response with some genes showing little or no differential expression and other genes showing a significant increase or decrease in gene expression in response to heat shock (Table 1; Fig. 2).

Cold Shock Results. A log phase culture growing in YEP/ dextrose medium at 37°C was split in half. One half of the culture remained at 37°C while the other half of the culture was shifted to 18°C. mRNA was isolated from both cultures 1 h after cold shock for comparison on microarrays. As expected,

two known cold shock genes (TIP1, TIR1) were expressed at a significantly higher level in the cold-shocked culture. Genes in other functional categories, such as glucose metabolism and heat shock displayed a mixed response with expression of some genes being unaffected and other genes exhibiting significant up- or down-regulation in response to cold shock (Table 2).

Steady-State Galactose vs. Glucose Results. mRNA was isolated from steady-state log phase YEP galactose and YEP glucose grown cultures for comparison on the microarrays. As expected, the GAL genes were expressed at a much higher level in the galactose culture. Many genes were differentially expressed in these cultures that were not a priori expected to exhibit differential expression. For example, some genes in the amino acid catabolic category were up-regulated in the galactose culture whereas genes in the one-carbon metabolism and purine categories were largely or entirely down-regulated in the galactose culture (Table 3). Genes in other categories, such as amino acid synthesis, abc transporter, cytochrome c, and cytochrome b, exhibited mixed responses; some genes in a category showed little or no obvious differential expression whereas other genes in the same category showed significant differential expression in the galactose and glucose cultures.

DISCUSSION

The results of these experiments show that many genes are differentially expressed under the three environmental conditions described here. The expected and predicted changes in gene expression, such as HSP12 in the heat-shocked culture, TIP1 in the cold-shocked culture, and GAL2 in the steady-state galactose culture, were observed in every case. However, in addition to the expected changes in gene expression, significant differential expression was also observed for many other genes that would not, a priori, be expected to be differentially expressed. For example, expression of PHO11 decreased and expression of YLR194, KIN2, and HXT6 increased in the heat shocked culture: Expression of MST1 and APE3 decreased and expression of PDR5 and GAR1 increased in the cold-shocked culture. In addition, ADE4 and SER2 were expressed at reduced levels whereas PHO84 and ACH1 were expressed at higher levels in cells grown in galactose compared with cells grown in glucose. Differential expression of these and many other genes was specific to one of these three environmental conditions.

. Many other genes were found to be differentially expressed under more than one condition. When differentially expressed genes in cold- and heat-shocked cultures were compared, 30 genes were found in common. Of these 30 genes, 28 showed inverse expression (i.e., increased expression under one condition and decreased expression under the other condition). Two genes, YCR058 and YKL102, showed elevated expression in response to both cold and heat shock. Fifteen genes were found to be differentially expressed in both the heat-shocked and steady-state galactose cultures: 9 genes showed increased expression and 5 showed decreased expression under both conditions. Twenty genes were differentially expressed in both the cold-shocked and steady-state galactose cultures: 8 genes showed decreased expression and 5 genes showed increased expression under both conditions. Six genes showed increased expression in the galactose culture and decreased expression in the cold shocked culture. One gene (ODP1) showed increased expression in both the cold-shocked and steady-state galactose cultures.

Gene expression is affected in a global fashion when environmental conditions are changed and both expected and unexpected genes are affected. There is also overlap in the genes that are differentially expressed under quite different environmental conditions. These results can be rationalized by considering the high degree of cross-pathway regulation in yeast. For example, there is evidence for cross-pathway regulation between (i) carbon and nitrogen metabolism (18), (ii) phosphate and sulfate metabolism (19), and (iii) purine, phosphate, and amino acid metabolism (20-24). There are also examples of the interaction of general and specific transcription factors (25, 26). Finally, within the broad class of amino acid biosynthetic genes, there is evidence for amino acid specific regulation of some genes, regulation via general control for other genes, and regulation via both specific and general control for other genes (22, 27-30).

Cross-pathway regulation arises from the complex structure of promoters. Virtually all promoters contain sites for multiple transcription factors and, therefore, virtually all genes are subject to combinatorial regulation. For example, the HIS4 promoter contains binding sites for GCN4 (the general amino acid control transcription factor), PHO2/BAS2 (a transcriptional regulator of phosphatase and purine biosynthetic genes), and BAS1 (a transcriptional regulator of purine biosynthetic genes) (31). It is likely that the complex effects on gene expression described in this work are a direct consequence of the combinatorial regulation of gene expression.

These findings illustrate the power of the highly parallel whole genome approach when examining gene expression. The global effects of environmental change on gene expression can now be directly visualized. It is clear that determining the mechanism(s) and the functional role of the dramatic global effects on gene

expression in different environments will be a significant challenge. The era of whole genome analysis will, ultimately, allow researchers to switch from the very focused single gene/promoter view of gene expression and instead view the cell more as a large complex network of gene regulatory pathways.

With the entire sequence of this model organism known, new approaches have been developed that allow for genome wide analyses (32, 33) of gene function. The genome microarrays represent a novel tool for genetic and expression analysis of the yeast genome. This pilot study uses arrays containing >35% of the yeast ORFs and it is clear that the entire set of ORFs from the yeast genome can be arrayed using the directed primer based strategy detailed here. Recent advances in arraying technology will allow all 6,100 ORFs to be arrayed in an area of less than 1.8 cm². Furthermore, as the technology improves, detection limits will allow less than 500 ng of starting mRNA material to be used for making probe.

The genome arrays provide for a robust, fully automated approach toward examining genome structure and gene function. They allow for comparisons between different genomes as well as a detailed study of gene expression at the global level. This research will help to elucidate relationships between genes and allow the researcher to understand gene function by understanding expression patterns across the yeast genome.

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- Fleischmann, R. D., Adams, M. D., White, O., Clayton, R. A., Kirkness, E. F., et al. (1995) Science 269, 496-512.

 Fraser, C. M., Gocsyne, J. D., White, O., Adams, M. D., Clayton, R. A., et al. (1995) Science 270, 397-403.

 Bult, C. J., White, O., Olsen, G. J., Zhou, L., Fleischmann, R. D., et al. (1996) Science 273, 1058-1073.

 Sulston, J. Du, Z., Thomas, K. Wilson, R. Millier, L. et al. (1992) Message
- 4. Sulston, J., Du, Z., Thomas, K., Wilson, R., Hillier, L., et al. (1992) Nature (London) 356, 37.
- Newman, T., de Brutjin, F. J., Green, P., Keegstra, K., Kende, H., et al. (1994) Plant Physiol. 106, 1241-1255.
- Physiol. 196, 1241-1255.
 Lashkari, D. A., Hunicke-Smith, S. P., Norgren, R. M., Davis, R. W. & Brennan, T. (1995) Proc. Nat. Acad. Sci. USA 92, 7912-7915.
 Schena, M., Shalon, D., Davis, R. W. & Brown, P. O. (1995) Science 270, 467-470.
 Shalon, D., Smith, S. & Brown, P. O. (1996) Genome Res. 6, 639-645.
 Heller, R. A., Schena, M., Chai, A., Shalon, D., Bedilion, T., Gilmore, J., Woolley, D. E. & Davis, R. W. (1997) Proc. Natl. Acad. Sci. USA 94, 2150-2155.
 DeRisi, J., Penland, L., brown, P. O., Bittner, M. L., Meltzer, P. S., Ray, M., Chen, Y.- Su Ya & Trent, J. M. (1996) Nat. Genet. 14, 457-460.
 Nelson, S. F., McCusker, J. H., Sander, M. A., Kee, Y., Modrich, P. & Brown P. O. (1993) Nat. Genet. 4, 11-18.
- (1993) Nat. Genet. 4, 11–18. Hoffman, C. S. & Winston, F. (1989) Gene 84, 473–479.
- Schmitt, M., Brown, T. & Trumpower, B. (1990) Nucleic Acids Res. 18, 3091. Ehrenhofer-Murray, A. E., Wurgler, F. E. & Sengstag, C. (1994) Mol. Gen. Genet.
- Kim, K-W., Kamerud, J. Q., Livingston, D. M. & Roon, R. J. (1988) J. Biol. Chem. 263, 11948–11953.
- Kim, K.-W. & Roon, R. J. (1984) J. Bacteriol. 157, 958-961.
 Craig, E. A. (1992) in The Molecular Biology of the Yeast Saccharomyces: Gene Expression, eds. Jones, E. W., Pringle, J. R. & Broach, J. R. (Cold Spring Harbor Lab, Press, Plainview, NY), Vol. 2, pp. 501-537.
 Dang, V. D., Bohn, C., Bolotin-Fukuhara, M. & Daignan-Fornier, B. (1996) J. Bacteriol. 178, 1842-1849.
- O'Connell, K. F. & Baker, R. E. (1992) Genetics 132, 63-73.

 Braus, G., Mosch, H. U., Vogel, K., Hinnen, A. & Hutter, R. (1989) EMBO J. 8,
- 939-933.
 Mosch, H. U., Scheier, B., Lahti, R., Mantsala, P. & Braus, G. H. (1991) J. Biol. Chem. 266, 20453-20456.
- Mitchell, A. P. & Magasanik, B. (1984) Mol. Cell. Biol. 4, 2767-2773.

 Daignan-Fornier, B. & Fink, G. R. (1992) Proc. Natl. Acad. Sci USA 89, 6746-6750.

- Tics-Baldwin, K., Fink, G. R. & Arndt, K. T. (1989) Science 246, 931-935.

 Mcssenguy, F. & Dubois, E. (1993) Mol. Cell. Biol. 13, 2586-2592.

 Devlin, C., Tice-Baldwin, K., Shore, D. & Arndt, K. T. (1991) Mol. Cell. Biol. 11,
- 3642-3651.

 Magasanik, B. (1992) in The Molecular and Cellular Biology of the Yeast Saccharomyces: Gene Expression. cds. Jones, E. W., Pringle, J. R. & Broach, J. R. (Cold Spring Harbor Lab. Press, Plainview, NY). Vol. 2, pp. 283-317.

 Hinnebusch, A. G. (1992) in The Molecular and Cellular Biology of the Yeast Saccharomyces: Gene Expression, cds. Jones, E. W., Pringle, J. R. & Broach, J. R. (Cold Spring Harbor Lab. Press, Plainview, NY), Vol. 2, pp. 319-414.

 Brisco, P. R. & Kohlhaw, G. B. (1990) J. Biol. Chem. 245, 11667-11675.

 O'Connell, K. F., Surdin-Kerjan, Y. & Baker R. E. (1995) Mol. Cell. Biol. 15, 1879-1888.

- Arndt K. T., Styles, C. & Fink, G. R. (1987) Science 237, 874-880. Smith, V., Chou, K. N., Lashkari, D., Botstein, D. & Brown, P. O. (1996) Science 274, 2069-2074.
- Shocmaker, D. D., Lashkari, D. A., Morris, D., Mittman, M. & Davis, R. W. (1996) Nat. Genet. 14, 450-456.

State Same

Fischer-Vze, Science 270, 1828 (1995).

35. T. C. James and S. C. Elgin, Mol. Cell Biol. 6, 3862 (1986); R. Paro and D. S. Hogness, Proc. Natl. Acad. Sci. U.S.A 88, 263 (1991); B. Tschiersch et al. EMBO J.: 13, 3822 (1994); M. T. Madireddi et al., Cell ... 87, 75 (1996); D. G. Stokes, K. D. Tartof, R. P. Perry, Proc. Natl. Acad. Sci. U.S.A. 93, 7137 (1996).

36. P. M. Patossari et al., J. Biol. Chem. 266, 10750 38. W. Hendriks et al., J. Cell Biochem. 59, 418 (1995)

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Grund, R. Sichenlaub, Appl. Environ. Microbiol. 58, ... 4068 (1992); V. Sharma, K. Suvarna, R. Meganathan, M. E. Hudspeth, J. Bacteriol. 174, 5057 (1992); M. Kanazawa et al., Enzyme Protein 47, 9 (1993); Z. L. Boynton, G. N. Bennet, F. B. Rudolph, -J. Bacteriol. 178, 3015 (1996).

37. M. Ho et al., Cell 77, 869 (1994).

(1991); A. Schmitz, K. H. Gartemann, J. Fiedler, E. 1939. We thank H. Skaletsky and F. Lewitter for help with

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Exploring the Metabolic and Genetic Control of Gene Expression on a Genomic Scale

Joseph L. DeRisi, Vishwanath R. Iyer, Patrick O. Brown*

DNA microarrays containing virtually every gene of Saccharomyces cerevisiae were used to carry out a comprehensive investigation of the temporal program of gene expression accompanying the metabolic shift from fermentation to respiration. The expression profiles observed for genes with known metabolic functions pointed to features of the metabolic reprogramming that occur during the diauxic shift, and the expression patterns of many previously uncharacterized genes provided clues to their possible functions. The same DNA microarrays were also used to identify genes whose expression was affected by deletion of the transcriptional co-repressor TUP1 or overexpression of the transcriptional activator YAP1. These results demonstrate the feasibility and utility of this approach to genomewide exploration of gene expression patterns.

The complete sequences of nearly a dozen microbial genomes are known, and in the next several years we expect to know the complete genome sequences of several metazoans, including the human genome. Defining the role of each gene in these genomes will be a formidable task, and understanding how the genome functions as a whole in the complex natural history of a living organism presents an even greater challenge.

Knowing when and where a gene is expressed often provides a strong clue as to its biological role. Conversely, the pattern of genes expressed in a cell can provide detailed information about its state. Although regulation of protein abundance in a cell is by no means accomplished solely by regulation of mRNA, virtually all differences in cell type or state are correlated with changes in the mRNA levels of many genes. This is fortuitous because the only specific reagent required to measure the abundance of the mRNA for a specific gene is a cDNA sequence. DNA microarrays, consisting of thousands of individual gene sequences printed in a high-density array on a glass microscope slide (1, 2), provide a practical and economical tool for studying gene expression on a very large scale (3-6).

Saccharomyces cerevisiae is an especially

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favorable organism in which to conduct a systematic investigation of gene expression. The genes are easy to recognize in the genome sequence, cis regulatory elements are generally compact and close to the transcription units, much is already known about its genetic regulatory mechanisms, and a powerful set of tools is available for its analysis.

A recurring cycle in the natural history of yeast involves a shift from anaerobic (fermentation) to aerobic (respiration) metabolism. Inoculation of yeast into a medium rich in sugar is followed by rapid growth fueled by fermentation, with the production of ethanol. When the fermentable sugar is exhausted, the yeast cells turn to ethanol as a carbon source for aerobic growth. This switch from anaerobic growth to aerobic respiration upon depletion of glucose, referred to as the diauxic shift, is correlated with widespread changes in the expression of genes involved in fundamental cellular processes such as carbon metabolism, protein synthesis, and carbohydrate storage (7). We used DNA microarrays to characterize the changes in gene expression that take place during this process for nearly the entire genome, and to investigate the genetic circuitry that regulates and executes this program.

Yeast open reading frames (ORFs) were amplified by the polymerase chain reaction (PCR), with a commercially available set of primer pairs (8). DNA microarrays, containing approximately 6400 distinct DNA sequences, were printed onto glass slides by

The Manual of Contract to using a simple robotic printing device (9). Cells from an exponentially growing culture of yeast were inoculated into fresh medium and grown at 30°C for 21 hours. After an initial 9 hours of growth, samples were harvested at seven successive 2-hour intervals, and mRNA was isolated (10). Fluorescently labeled cDNA was prepared by reverse transcription in the presence of Cy3(green)or Cy5(red)-labeled deoxyuridine triphosphate (dUTP) (11) and then hybridized to the microarrays (12). To maximize the reliability with which changes in expression levels could be discerned, we labeled cDNA prepared from cells at each successive time. point with Cy5, then mixed it with a Cy3labeled "reference" cDNA sample prepared from cells harvested at the first interval after inoculation. In this experimental design, the relative fluorescence intensity measured for the Cy3 and Cy5 fluors at each array element provides a reliable measure of the relative abundance of the corresponding mRNA in the two cell populations (Fig. 1). Data from the series of seven samples (Fig. 2), consisting of more than 43,000 expression-ratio measurements, were organized into a database to facilitate efficient exploration and analysis of the results. This database is publicly available on the Internet (13).

During exponential growth in glucoserich medium, the global pattern of gene expression was remarkably stable. Indeed, when gene expression patterns between the first two cell samples (harvested at a 2-hour interval) were compared, mRNA levels differed by a factor of 2 or more for only 19 genes (0.3%), and the largest of these differences was only 2.7-fold (14). However, as glucose was progressively depleted from the growth media during the course of the experiment, a marked change was seen in the global pattern of gene expression. mRNA levels for approximately 710 genes were induced by a factor of at least 2, and the mRNA levels for approximately 1030 genes declined by a factor of at least 2. Messenger RNA levels for 183 genes increased by a factor of at least 4, and mRNA levels for 203 genes diminished by a factor of at least 4. About half of these differentially expressed genes have no currently recognized function and are not yet named. Indeed, more than 400 of the differentially expressed genes have no apparent homology

to any gene whose function is known (15). The responses of these previously uncharacterized genes to the diauxic shift therefore provides the first small clue to their possible roles.

The global view of changes in expression of genes with known functions provides a vivid picture of the way in which the cell adapts to a changing environment. Figure 3 shows a portion of the yeast metabolic pathways involved in carbon and energy metabolism. Mapping the changes we observed in the mRNAs encoding each enzyme onto this framework allowed us to infer the redirection in the flow of metabolites through this system. We observed large inductions of the genes coding for the enzymes aldehyde dehydrogenase (ALD2) and acetyl-coenzyme A(CoA) synthase (ACS1), which function together to convert the products of alcohol dehydrogenase into acetyl-CoA, which in turn is used to fuel the tricarboxylic acid (TCA) cycle and the glyoxylate cycle. The concomitant shutdown of transcription of the genes encoding pyruvate decarboxylase and induction of pyruvate carboxylase rechannels pyruvate away from acetaldehyde, and instead to oxalacetate, where it can serve to supply the TCA cycle and gluconeogenesis. Induction of the pivotal genes PCK1, encoding phosphoenolpyruvate carboxykinase, and FBP1, encoding fructose 1,6-biphosphatase, switches the directions of two key irreversible steps in glycolysis, reversing the flow of metabolites along the reversible steps of the glycolytic pathway toward the essential biosynthetic precursor, glucose-6-phosphate. Induction of the genes coding for the trehalose synthase and glycogen synthase complexes promotes channeling of glucose-6-phosphate into these carbohydrate storage pathways.

Just as the changes in expression of genes encoding pivotal enzymes can provide insight into metabolic reprogramming, the behavior of large groups of functionally related genes can provide a broad view of the systematic way in which the yeast cell adapts to a changing environment (Fig. 4). Several classes of genes, such as cytochrome c-related genes and those involved in the TCA/glyoxylate cycle and carbohydrate storage, were coordinately induced by glucose exhaustion. In contrast, genes devoted to protein synthesis, including ribosomal proteins, tRNA synthetases, and translation, elongation, and initiation factors, exhibited a coordinated decrease in expression. More than 95% of ribosomal genes showed at least twofold decreases in expression during the diauxic shift (Fig. 4) (13). A noteworthy and illuminating exception was that the

genes encoding mitochondrial ribosomal genes were generally induced rather than repressed after glucose limitation, highlighting the requirement for mitchondrial biogenesis (13). As more is learned about the functions of every gene in the yeast genome, the ability to gain insight into a cell's response to a changing environment through its global gene expression patterns will become increasingly powerful.

Several distinct temporal patterns of expression could be recognized, and sets of genes could be grouped on the basis of the similarities in their expression patterns. The characterized members of each of these groups also shared important similarities in their functions. Moreover, in most cases, common regulatory mechanisms could be inferred for sets of genes with similar expression profiles. For example, seven genes showed a late induction profile, with mRNA levels increasing by more than ninefold at

the last timepoint but less than threefold at the preceding timepoint (Fig. 5B). All of these genes were known to be glucose-repressed, and five of the seven were previously noted to share a common upstream activating sequence (UAS), the carbon source response element (CSRE) (16-20). A search in the promoter regions of the remaining two genes, ACRI and IDP2, revealed that ACRI, a gene essential for ACSI activity, also possessed a consensus CSRE motif, but interestingly, IDP2 did not. A search of the entire yeast genome sequence for the consensus CSRE motif revealed only four additional candidate genes, none of which showed a similar induction.

Examples from additional groups of genes that shared expression profiles are illustrated in Fig. 5, C through F. The sequences upstream of the named genes in Fig. 5C all contain stress response elements (STRE), and with the exception

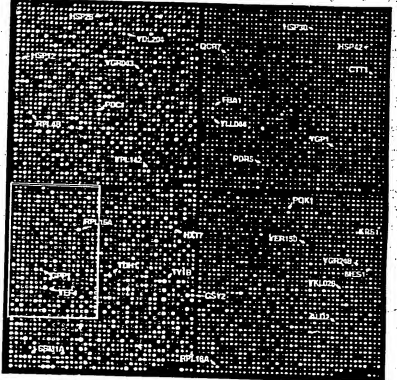


Fig. 1. Yeast genome microarray. The actual size of the microarray is 18 mm by 18 mm. The microarray was printed as described (9). This image was obtained with the same fluorescent scanning confocal microscope used to collect all the data we report (49). A fluorescently labeled cDNA probe was prepared from mRNA isolated from cells harvested shortly after inoculation (culture density of <5 × 10° cells/ml and media glucose level of 19 g/liter) by reverse transcription in the presence of Cy3-dUTP. Similarly, a second probe was prepared from mRNA isolated from cells taken from the same culture 9.5 hours later (culture density of ~2 × 10° cells/ml, with a glucose level of <0.2 g/liter) by reverse transcription in the presence of Cy5-dUTP. In this image, hybridization of the Cy3-dUTP-labeled cDNA (that is, mRNA expression at the initial timepoint) is represented as a green signal, and hybridization of Cy5-dUTP-labeled cDNA (that is, mRNA expression at 9.5 hours) is represented as a red signal. Thus, genes induced or repressed after the diauxic shift appear in this image as yellow spots.

of HSP42, have previously been shown to be controlled at least in part by these elements (21-24). Inspection of the sequences upstream of HSP42 and the two uncharacterized genes shown in Fig. 5C, YKL026c, a hypothetical protein with similarity to glutathione peroxidase, and YGR043c, a putative transaldolase, revealed that each of these genes also possess repeated upstream copies of the stressresponsive CCCCT motif. Of the 13 additional genes in the yeast genome that shared this expression profile [including HSP30, ALD2, OM45, and 10 uncharacterized ORFs (25)], nine contained one or more recognizable STRE sites in their upstream regions.

The heterotrimeric transcriptional activator complex HAP2,3,4 has been shown to be responsible for induction of several genes important for respiration (26-28). This complex binds a degenerate consensus sequence known as the CCAAT box (26). Computer analysis, using the consensus sequence TNRYTGGB (29), has suggested that a large number of genes involved in respiration may be specific targets of HAP2,3,4 (30). Indeed, a putative HAP2,3,4 binding site could be found in the sequences upstream of each of the seven cytochrome c-related genes that showed the greatest magnitude of induction (Fig. 5D). Of 12 additional cytochrome c-related genes that were induced, HAP2,3,4 binding sites were present in all but one. Significantly, we found that transcription of HAP4 itself was induced nearly ninefold concomitant with the diauxic shift.

Control of ribosomal protein biogenesis is mainly exerted at the transcriptional level, through the presence of a common upstream-activating element (UAS_{rpg}) that is recognized by the Rap1 DNA-binding protein (31, 32). The expression profiles of seven ribosomal proteins are shown in Fig. 5F. A search of the sequences upstream of all seven genes revealed consensus Rap1-binding motifs (33). It has been suggested that declining Rap1 levels in the cell during starvation may be responsible for the decline in ribosomal protein gene expression (34). Indeed, we observed that the abundance of RAPI mRNA diminished by 4.4-fold, at about the time of glucose exhaustion.

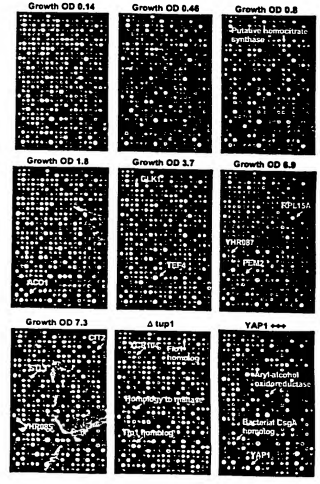
Of the 149 genes that encode known or putative transcription factors, only two, HAP4 and SIP4, were induced by a factor of more than threefold at the diauxic shift. SIP4 encodes a DNA-binding transcriptional activator that has been shown to interact with Snf1, the "master regulator" of glucose repression (35). The eightfold induction of SIP4 upon depletion of glucose strongly suggests a role in the induction of

of HSP42, have previously been shown to downstream genes at the diauxic shift.

Although most of the transcriptional responses that we observed were not previously known, the responses of many genes during the diauxic shift have been described. Comparison of the results we obtained by DNA microarray hybridization with previously reported results therefore provided a strong test of the sensitivity and accuracy of this approach. The expression patterns we observed for previously characterized genes showed almost perfect concordance with previously published results (36). Moreover, the differential expression measurements obtained by DNA microarray hybridization were reproducible in duplicate experiments. For example, the remarkable changes in gene expression between cells harvested immediately after inoculation and immediately after the diauxic shift (the first and sixth intervals in this time series) were measured in duplicate, independent DNA microarray hybridizations. The correlation coefficient for two complete sets of expression ratio measurements was 0.87, and for more than 95% of the genes, the expression ratios measured in these duplicate experiments differed by less than a factor of 2. However, in a few cases, there were discrepancies between our results and previous results, pointing to technical limitations that will need to be addressed as DNA microarray technology advances (37, 38). Despite the noted exceptions, the high concordance between the results we obtained in these experiments and those of previous studies provides confidence in the reliability and thoroughness of the survey.

The changes in gene expression during this diauxic shift are complex and involve integration of many kinds of information about the nutritional and metabolic state of the cell. The large number of genes whose expression is altered and the diversity of temporal expression profiles observed in this experiment highlight the challenge of understanding the underlying regulatory mechanisms. One approach to defining the contributions of individual regulatory genes to a complex program of this kind is to use DNA microarrays to identify genes whose expression is affected

Fig. 2. The section of the array indicated by the gray box in Fig. 1 is shown for each of the experiments described here. Representative genes are labeled. In each of the arrays used to analyze gene expression during the diauxic shift, red spots represent genes that were induced relative to the initial timepoint. and green spots represent genes that were repressed relative to the initial timepoint. In the arrays used to analyze the effects of the tup14 mutation and YAP1 overexpression, red spots represent genes whose expression was increased, and green spots represent genes whose expression was decreased by the genetic modification. Note that distinct sets of genes are induced and repressed in the different experiments. The complete images of each of these arrays can be viewed on the Internet (13). Cell density as measured by optical density (OD) at 600 nm was used to measure the growth of the



by mutations in each putative regulatory gene. As a test of this strategy, we analyzed the genomewide changes in gene expression that result from deletion of the TUPI gene. Transcriptional repression of many genes by glucose requires the DNA-binding repressor

Migl and is mediated by recruiting the transcriptional co-repressors Tupl and Cyc8/ Ssn6 (39). Tup1 has also been implicated in repression of oxygen-regulated, mating-typespecific, and DNA-damage-inducible genes (40). 'X X

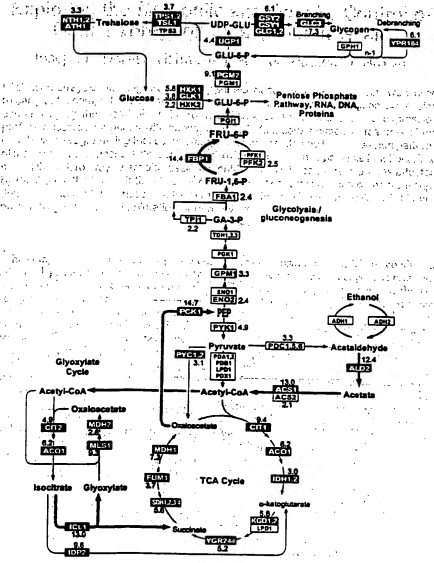


Fig. 3. Metabolic reprogramming inferred from global analysis of changes in gene expression. Only key metabolic intermediates are identified. The yeast genes encoding the enzymes that catalyze each step in this metabolic circuit are identified by name in the boxes. The genes encoding succinyl-CoA synthase and glycogen-debranching enzyme have not been explicitly identified, but the ORFs YGR244 and YPR184 show significant homology to known succinyl-CoA synthase and glycogen-debranching enzymes, respectively, and are therefore included in the corresponding steps in this figure. Red boxes with white lettering identify genes whose expression increases in the diauxic shift. Green boxes with dark green lettering identify genes whose expression diminishes in the diauxic shift. The magnitude of induction or repression is indicated for these genes. For multimeric enzyme complexes, such as succinate dehydrogenase, the indicated fold-induction represents an unweighted average of all the genes listed in the box. Black and white boxes indicate no significant differential expression (less than twofold). The direction of the arrows connecting reversible enzymatic steps indicate the direction of the flow of metabolic intermediates, inferred from the gene expression pattern, after the diauxic shift. Arrows representing steps catalyzed by genes whose expression was strongly induced are highlighted in red. The broad gray arrows represent major increases in the flow of metabolites after the diauxic shift, inferred from the indicated changes in gene expression.

Wild-type yeast cells and cells bearing a deletion of the TUP1 gene (mp1 \D) were grown in parallel cultures in rich medium containing glucose as the carbon source. Messenger RNA was isolated from exponentially growing cells from the two populations and used to prepare cDNA labeled with Cy3 (green) and Cy5 (red), respectively (11). The labeled probes were mixed and simultaneously hybridized to the microarray. Red spots on the microarcray therefore represented genes whose transcription was induced in the nipl A strain, and thus presumably repressed by Tupl (41). A representative section of the microarray (Fig. 2, bottom middle panel) illustrates that the genes whose expression was affected by the tup 1 A mutation, were, in general, distinct from those induced upon glucose exhaustion [complete images of all the arrays shown in Fig. 2 are available on the Internet (13)]. Nevertheless, :34 (10%) of the genes that were induced by a factor of at least 2 after the diauxic shift were similarly induced by deletion of TUP1, suggesting that these genes may be subject to TUPI-mediated repression by glucose. For example, SUC2, the gene encoding invertase, and all five hexose transporter genes that were induced during the course of the diauxic shift were similarly induced, in duplicate experiments, by the deletion of TUP1.

The set of genes affected by Tupl in this experiment also included a-glucosidases, the mating-type-specific genes MFA1 and MFA2, and the DNA damage-inducible RNR2 and RNR4, as well as genes involved in flocculation and many genes of unknown function. The hybridization signal corresponding to expression of TUP1 itself was also severely reduced because of the (incomplete) deletion of the transcription unit in the $n\phi l\Delta$ strain, providing a positive control in the experiment (42).

Many of the transcriptional targets of Tupl fell into sets of genes with related biochemical functions. For instance, although only about 3% of all yeast genes appeared to be TUP1-repressed by a factor of more than 2 in duplicate experiments under these conditions, 6 of the 13 genes that have been implicated in flocculation (15) showed a reproducible increase in expression of at least twofold when TUP1 was deleted. Another group of related genes that appeared to be subject to TUPI repression encodes the serine-rich cell wall mannoproteins, such as Tip1 and Tirl/Srp1 which are induced by cold shock and other stresses (43), and similar, serine-poor proteins, the seripauperins (44). Messenger RNA levels for 23 of the 26 genes in this group were reproducibly elevated by at least 2.5-fold in the tup 1 A

strain, and 18 of these genes were induced by more than sevenfold when TUP1 was deleted. In contrast, none of 83 genes that could be classified as putative regulators of the cell division cycle were induced more than twofold by deletion of TUP1. Thus, despite the diversity of the regulatory systems that employ Tup1, most of the genes that it regulates under these conditions fall into a limited number of distinct functional classes.

Because the microarray allows us to monitor expression of nearly every gene in yeast, we can, in principle, use this approach to identify all the transcriptional targets of a regulatory protein like Tup1. It is important to note, however, that in any single experiment of this kind we can only recognize those target genes that are normally repressed (or induced) under the conditions of the experiment. For instance, the experiment described here analyzed a MAT a strain in which MFAI and MFA2, the genes encoding the afactor mating pheromone precursor, are normally repressed. In the isogenic tup! strain, these genes were inappropriately expressed, reflecting the role that Tup1 plays in their repression. Had we instead carried out this experiment with a MATA strain (in which expression of MFA1 and MFA2 is not repressed), it would not have been possible to conclude anything regarding the role of Tupl in the repression of these genes. Conversely, we cannot distinguish indirect effects of the chronic absence of Tup1 in the mutant strain from effects directly attributable to its participation in repressing the transcription of a

Another simple route to modulating the activity of a regulatory factor is to overexpress the gene that encodes it. YAPI encodes a DNA-binding transcription factor belonging to the b-zip class of DNA-binding proteins. Overexpression of YAP1 in yeast confers increased resistance to hydrogen peroxide, o-phenanthroline, heavy metals, and osmotic stress (45). We analyzed differential gene expression between a wild-type strain bearing a control plasmid and a strain with a plasmid expressing YAP1 under the control of the strong GAL1-10 promoter, both grown in galactose (that is, a condition that induces YAP1 overexpression). Complementary DNA from the control and YAP1 overexpressing strains, labeled with Cy3 and Cy5, respectively, was prepared from mRNA isolated from the two strains and hybridized to the microarray. Thus, red spots on the array represent genes that were induced in the strain overexpressing YAPI.

Of the 17 genes whose mRNA levels increased by more than threefold when

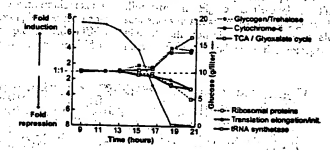
YAPI was overexpressed in this way, five bear homology to aryl-alcohol oxidoreductases (Fig. 2 and Table 1). An additional four of the genes in this set also belong to the general class of dehydrogenases/oxidoreductases. Very little is known about the role of aryl-alcohol oxidoreductases in S. cerevisiae, but these enzymes have been isolated from ligninolytic fungi, in which they participate in coupled redox reactions, oxidizing aromatic, and aliphatic unsaturated alcohols to aldehydes with the production of hydrogen peroxide (46, 47). The fact that a remarkable fraction of the targets identified in this experiment belong to the same small, functional group of oxidoreductases suggests that these genes

might play an important protective role during oxidative stress. Transcription of a small number of genes was reduced in the strain overexpressing Yapl. Interestingly many of these genes encode sugar permeases or enzymes involved in inositol metabolism.

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We searched for Yapl-binding sites (TTACTAA or TGACTAA) in the sequences upstream of the target genes we identified (48). About two-thirds of the genes that were induced by more than threefold upon Yapl overexpression had one or more binding sites within 600 bases upstream of the start codon (Table 1), suggesting that they are directly regulated by Yapl. The absence of canonical Yapl-bind-

Fig. 4. Coordinated regulation of functionally related genes. The curves represent the average induction or repression ratios for all the genes in each indicated group. The total number of genes in each group was as follows: ribosomal proteins, 112; translation elongation and initiation



factors, 25; tRNA synthetases (excluding mitochondial synthetases), 17; glycogen and trehalose synthesis and degradation, 15; cytochrome c oxidase and reductase proteins, 19; and TCA- and glyoxylate-cycle enzymes, 24.

Table 1. Genes induced by YAP1 overexpression. This list includes all the genes for which mRNA levels increased by more than twofold upon YAP1 overexpression in both of two duplicate experiments, and for which the average increase in mRNA level in the two experiments was greater than threefold (50). Positions of the canonical Yap1 binding sites upstream of the start codon, when present, and the average fold-increase in mRNA levels measured in the two experiments are indicated.

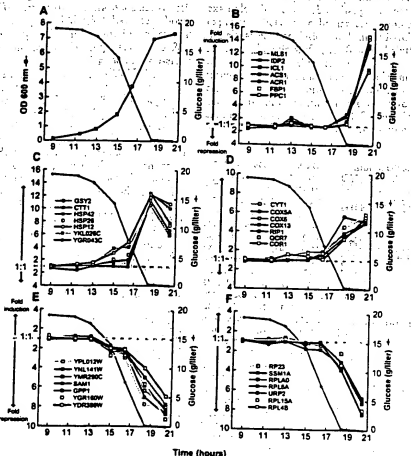
ORF	Distance of Yap1 site from ATG	Gene	Description	Fold- increase
YNL331C			Putative aryl-alcohol reductase	ν
YKL071W	162-222 (5 sites)		Similarity to bacterial csgA protein	12.9
YML007W		YAP1	Transcriptional activator involved in oxidative stress response	10.4 9.8
YFL056C	223, 242	•	Homology to anyl-alcohol dehydrogenases	9.0
YLL060C :	- 98 ·		Putative glutathione transferase	7.4
YOL165C	266		Putative aryl-alcohol dehydrogenase (NADP+)	7.0
YCR107W			Putative aryl-alcohol reductase	6.5
YML116W	409	ATR1	Aminotriazole and 4-nitroquinoline resistance protein	6.5
YBR008C	142, 167, 364		Homology to benomy/methotrexate resistance protein	6.1
YCLX08C			Hypothetical protein	
YJR155W			Putative aryl-alcohol dehydrogenase	6.1
YPL171C	148, 212	OYE3	NAPDH dehydrogenase (old yellow enzyme), isoform 3	6.0 5.8
YLR460C	167, 317		Homology to hypothetical proteins YCR102c and YNL134c	4.7
YKR076W	178		Homology to hypothetical protein YMR251w	4.5
YHR179W	327	OYE2	NAD(P)H oxidoreductase (old yellow enzyme), isoform 1	4.1
YML131W	507		Similarity to A. thaliana zeta-crystallin homolog	3.7
YOL126C		MDH2	Malate dehydrogenase	3.3

ing sites upstream of the others may reflect an ability of Yapl to bind sites that differ from the canonical binding sites, perhaps in cooperation with other factors, or less likely, may represent an indirect effect of Yap1 overexpression, mediated by one or more intermediary factors. Yapl sites were found only four times in the corresponding region of an arbitrary set of 30 genes that were not differentially regulated by Yap1.

Use of a DNA microarray to characterize the transcriptional consequences of mutations affecting the activity of regulatory molecules provides a simple and powerful approach to dissection and characterization of regulatory pathways and net-

works. This strategy also has an important a required for fabricating and using DNA practical application in drug screening. Mutations in specific genes encoding candidate drug targets can serve as surrogates ofor the ideal chemical inhibitor or modulator of their activity. DNA microarrays can be used to define the resulting signature pattern of alterations in gene expresasion, and then subsequently used in an assay to screen for compounds that reproduce the desired signature pattern.

DNA microarrays provide a simple and economical way to explore gene expression patterns on a genomic scale. The hurdles to extending this approach to any other organism are minor. The equipment



Time (hours) Fig. 5. Distinct temporal patterns of induction or repression help to group genes that share regulatory properties. (A) Temporal profile of the cell density, as measured by OD at 600 nm and glucose concentration in the media. (B) Seven genes exhibited a strong induction (greater than ninefold) only at the last timepoint (20.5 hours). With the exception of IDP2, each of these genes has a CSRE UAS. There were no additional genes observed to match this profile. (C) Seven members of a class of genes marked by early induction with a peak in mRNA levels at 18.5 hours. Each of these genes contain STRE motif repeats in their upstream promoter regions. (D) Cytochrome c oxidase and ubiquinol cytochrome c reductase genes. Marked by an induction coincident with the diauxic shift, each of these genes contains a consensus binding motif for the HAP2,3,4 protein complex. At least 17 genes shared a similar expression profile. (E) SAM1, GPP1, and several genes of unknown function are repressed before the diauxic shift, and continue to be repressed upon entry into stationary phase. (F) Ribosomal protein genes comprise a large class of genes that are repressed upon depletion of glucose. Each of the genes profiled here contains one or more RAP1-binding motifs upstream of its promoter. RAP1 is a transcriptional regulator of most ribosomal proteins.

emicroarrays (9) consists of components that were chosen for their modest cost and simplicity. It was feasible for a small group to accomplish the amplification of more than 6000 genes in about 4 months and, once the amplified gene sequences were in hand, only 2 days were required to print a set of 110 microarrays of 6400 elements each. Probe preparation, hybridization, and fluorescent imaging care also simple procedures. Even conceptually simple exoperiments, as we described here, can yield vast amounts of information. The value of the information from each experiment of this kind will progressively increase as more is learned about the functions of each gene and as additional experiments define the global changes in gene expression in diverse other natural processes and genetic perturbations. Perhaps the greatest challenge, now is to develop efficient methods for organizing, distributing, interpreting, and extracting insights from the large volumes of data these experiments will provide. There was to Father the

REFERENCES AND NOTES

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- 1. M. Schena, D. Shalon, R. W. Davis, P. O. Brown, Science 270, 467 (1995).
- 2. D. Shaton, S. J. Smith, P. O. Brown, Genome Res. 6, 639 (1996).
- 3. D. Lashkari, Proc. Natl. Acad. Sci. U.S.A.; in press. J. DeRisi et al., Nature Genet. 14, 457 (1996).
- 5. D. J. Lockhart et al., Nature Biotechnol. 14, 1675
- M. Chee et al., Science 274, 610 (1996).
- 7: M. Johnston and M. Carlson, in The Molecular Biology of the Yeast Saccharomyces: Gene Expres E. W. Jones, J. R. Pringle, J. R. Broach, Eds. (Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, 1992), p. 193.
- 8. Primers for each known or predicted protein coding sequence were supplied by Research Gener PCR was performed with the protocol supplied by Research Genetics, using genomic DNA from ye strain S288C as a template. Each PCR product was verified by agarose gel electrophoresis and we deemed correct if the lane contained a single band of appropriate mobility. Faitures were marked as such in the database. The overall success rate for a singlepass amplification of 6116 ORFs was ~94.5%.
- 9. Glass slides (Gold Seal) were cleaned for 2 hours in a solution of 2 N NaOH and 70% ethanol, After ring in distilled water, the slides were then treated with a 1:5 dilution of poly-L-lysine adhesive solution (Sigma) for 1 hour, and then dried for 5 min at 40°C in a vacuum oven, DNA samples from 100-µI PCR reactions were purified by ethanol purification in 96-well microtiter plates. The resulting precipitates were resuspended in 3x standard saline citrate (SSC) and transferred to new plates for arraying. A custom arraying robot was used to print on a batch of 110 slides. Details of the design of the microarrayer are available at crigm.stanford.edu/pbrown. After printing, the microarrays were rehydrated for 30 s in a humid chamber and then snap-dried for 2 s on a hot plate (100°C). The DNA was then ultraviolet (UV)crossinked to the surface by subjecting the slides to 60 mJ of energy (Stratagene Stratalinker). The rest of the poly-L-lysine surface was blocked by a 15-min incubation in a solution of 70 mM succinic anhydride dissolved in a solution consisting of 315 ml of 1-methyl-2-pyrrolidinone (Aldrich) and 35 ml of 1 M boric acid (pH 8.0). Directly after the blocking reac-

- tion, the bound DNA was denatured by a 2-min incubation in distilled water at ~95°C. The sides were then transferred into a bath of 100% ethanol at room temperature, rinsed; and then spun dry in a clinical centifluge. Sides were stored in a closed box at room temperature until used.
- 10. YPD medium (8 liters), in a 10-liter fermentation vessel, was inoculated with 2 ml of a fresh overnight culture of yeast strain DBY7286 (MATa, ura3, GAL2). The fermentor was maintained at 30°C with constant agitation and aeration. The glucose content of the media was measured with a UV test ldt (Boehringer Mannheim, catalog number 716251) Cell density was measured by OD at 600-nm wavelength. Aliquots of culture were rapidly withdrawn from the fermentation vessel by peristatic pump, spun down at room temperature, and then flash frozen with liquid nitrogen. Frozen cells were stored at -80°C.
- 11. Cy3-dUTP or Cy5-dUTP (Amersham) was incorporated during reverse transcription of 1.25 ag of polyadenylated [poly(A)+] RNA, primed by a dT(16) oligomer. This mixture was heated to 70°C for 10 min, and then transferred to ice. A premixed solution, consisting of 200 U Superscript II (Gibco), buffer, deoxyribonucleoside triphosphates, and fluorescent nucleotides, was added to the RNA. Nudectides were used at these final concentrations: 500 μM for dATP, dCTP, and dGTP and 200 μM for dTTP. Cy3-dUTP and Cy5-dUTP were used at a final concentration of 100 µM. The reaction was then incubated at 42°C for 2 hours. Unincorporated fluorescent nucleotides were removed by first diluting the reaction mixture with of 470 µJ of 10 mM tris-HCI (pH 8.0)/1 mM EDTA and then subsequently concentrating the mix to ~5 µl, using Centricon-30 microconcentrators (Amicon).
- 12. Purified, labeled cDNA was resuspended in 11 μl of 3.5× SSC containing 10 μg poly(dA) and 0.3 μl of 10% SDS. Before hybridization, the solution was boiled for 2 min and then allowed to cool to room temperature. The solution was applied to the microarray under a cover slip, and the slide was placed in a custom hybridization chamber which was subsequently incubated for ~8 to 12 hours in a water bath at 62°C. Before scanning, slides were washed in 2× SSC, 0.2% SDS for 5 min, and then 0.05× SSC for 1 min. Slides were dried before scanning by cantrifugation at 500 rpm in a Beckman CS-6R centrifuge.
- The complete data set is available on the internet at cmgm.stanford.edu/pbrown/explore/index.html
- 14. For 95% of all the genes analyzed, the mRNA levels measured in cells harvested at the first and second interval after inoculation differed by a factor of less than 1.5. The correlation coefficient for the comparison between mRNA levels measured for each gene in these two different mRNA samples was 0.98. When duplicate mRNA preparations from the same cell sample iwere compared in the same way, the correlation coefficient between the expression levels measured for the two samples by comparative hybridization was 0.99.
- 15. The numbers and identities of known and putative genes, and their homologies to other genes, were gathered from the following public databases: Seccharomycas Genome Database (genome-www, stanford.edu), Yeast Protein Database (quest?, proteome.com), and Munich Information Centre for Protein Sequences (speedy.mips.biochem.mpg.de/ mips/yeast/index.htmb/.
- A. Scholer and H. J. Schuller, Mol. Cell. Biol. 14, 3613 (1994).
- S. Kratzer and H. J. Schuller, Gene 161, 75 (1995).
 R. J. Haselbeck and H. L. McAlister, J. Biol. Chem.
- 268, 12116 (1983).
 M. Fernandez, E. Fernandez, R. Rodicio, Mal. Gen. Genet. 242, 727 (1994).
- 20. A. Hartig et al., Nucleic Acids Res. 20, 5677 (1992).
- 21. P. M. Martinez et al., EMBO J. 15, 2227 (1996).
- J. C. Varela, U. M. Praekett, P. A. Meacock, R. J. Planta, W. H. Mager, Mol. Cell. Biol. 15, 6232 (1995).
- H. Ruis and C. Schuller, Bioessays 17, 959 (1995).
 J. L. Parrou, M. A. Teste, J. Francois, Microbiology 143, 1891 (1997).

 This expression profile was defined as having an induction of greater than 10-fold at 18.5 hours and less than 11-fold at 20.5 hours.

- S. L. Forsburg and L. Guarente, Genes Dev. 3, 1166.
 (1989).
- J. T. Olesen and L. Guerente, bid. 4, 1714 (1990).
 M. Rosenkrantz, C. S. Kell, E. A. Pennet, L. J. Devenish, Mol. Microbiol. 13, 119 (1994).
- 29. Single-letter abbreviations for the amino acid residues are as follows: A, Ala; C, Cys; D, Asp; E, Glu; F, Phe; G, Gly; H, His; I, Ile; K, Lys; L, Leu; M, Met; N, Asn; P, Pro; O, Gin; R, Arg; S, Ser; T, Thr; V, Vaj; W, Trp; and Y, Tyr. The nucleotide codes are as follows: B-C, G, or T; N-G; A, T, or C; R-A or G; and Y-C or
- 30. C. Fondrat and A. Kalogeropoulos, Comput. Appl. Biosci. 12, 363 (1996).
- 31. D. Shore, Trends Genet. 10, 408 (1994).
- 32. R. J. Planta and H. A. Raue, ibid. 4, 64 (1988).
- 33. The degenerate consensus sequence VYCYRNNC-MNH was used to search for potential RAP1-binding sites. The exact consensus, as defined by (30), is WACAYCCRTACATYW, with up to three differences allowed.
- S. F. Neuman, S. Bhattacharya, J. R. Broach, Mol. Cell. Biol. 15, 3187 (1995).
- 35. P. Lesage, X. Yang, M. Cartson, ibid. 16, 1921 (1996).
- For example, we observed large inductions of the genes coding for PCK1, FBP1 [Z. Yin et al., Mol. Microbiol. 20, 751 (1996)], the central glyoxylate cycle gene ICL1 (A. Scholer and H. J. Schuller, Curr. Genet. 23, 375 (1993)), and the "aerobic" isoform of acetyl-CoA synthase, ACS1 [M. A. van den Berg et al., J. Biol. Chem. 271, 28953 (1996)]. with concomitant down-regulation of the glycolytic-specific genes PYK1 and PFK2 [P. A. Moore et al., Mol. Cell. Biol. 11, 5330 (1991)]. Other genes not directly involved in carbon metabolism but known to be induced upon nutrient limitation include genes encoding cytosolic catalase T CTT1 [P. H. Bissinger et al., ibid. 9, 1309 (1989)] and several genes encoding small heat-shock proteins, such as HSP12, HSP26, and HSP42 [I. Farkas et al., J. Biol. Chem. 266, 15602 (1991); U. M. Praekelt and P. A. Meacock, Mol. Gen. Genet. 223, 97 (1990); D. Wotton et al., J. Biol. Chem. 271, 2717 (1996)].
- 37. The levels of induction we measured for genes that were expressed at very low levels in the uninduced state (notably, FBP1 and PCK1) were generally lower than those previously reported. This discrepancy was likely due to the conservative background subtraction method we used, which generally resulted in overestimation of very low expression levels (46).
- 38. Cross-hybridization of highly related sequences can also occasionally obscure changes in gene expression, an important concern where members of gene families are functionally specialized and differentially regulated. The major alcohol dehydrogenase genes, ADH1 and ADH2, share 88% nucleotide identity. Reciprocal regulation of these genes is an important feature of the diauxic shift, but was not observed in this experiment, presumably because of cross-hybridization of the fluorescent cDNAs representing these two genes. Nevertheless, we were able to detect differential expression of closely related isoforms of other enzymes, such as HXK1/HXK2 (77% identical) [P. Herrero et al., Yeast 11, 137 (1995)], MLS1/ DAL7 (73% identical) (20), and PGM1/PGM2 (72% identical) [D. Oh, J. E. Hopper, Mol. Cell. Biol. 10, 1415 (1990)), in accord with previous studies. Use in the microarray of deliberately selected DNA sequences corresponding to the most divergent segments of homologous genes, in lieu of the complete gene sequences, should relieve this problem in many
- F. E. Williams, U. Varanasi, R. J. Trumbly, Mol. Cell. Biol. 11, 3307 (1991).
- D. Tzamarias and K. Struhl, Nature 369, 758 (1994).
 Differences in mRNA levels between the tup1 Δ and wild-type strain were measured in two independent experiments. The correlation coefficient between the complete sets of expression ratios measured in these duplicate experiments was 0.83. The concor-

- dance between the sets of genes that appeared to be induced was very high between the two experiments. When only, the 355 genes that showed at least a twofold increase in mRNA in the tup 14 strain in either of the duplicate experiments were compared, the complation coefficient was 0.82.
- 42. The tup14 mutation consists of an invention of the LEU2 coding sequence, including a stop codon, between the ATG of TUP1 and an Eco R I site 124 base pairs before the stop codon of the TUP1 gens.
- 43. L.R. Kowaiski, K. Kondo, M. Induye, Mcl. Microbiol. 15, 341 (1995).
- 44. M. Viswanathan, G. Muthulomer, Y.-S. Cong. J. Lenard, Gene 148, 149 (1994).
- 45. D. Hirata, K. Yano, T. Miyakawa, Mol. Gen. Genet. 242, 250 (1994).
- A. Gutierrez, L. Caramelo, A. Prieto, M. J. Martinez, A. T. Martinez, Appl. Environ. Microbiol. 60, 1783. (1994).
- A. Muheim et al., Eur. J. Biochem. 195, 369 (1991).
 J. A. Wemmie, M. S. Szczypka, D. J. Thiele, W. S. Moye-Rowley, J. Biol. Chem. 269, 32592 (1994).
- 49. Microarrays were scanned using a custom-built scanning laser microscope built by S. Smith with software written by N. Ziv. Details concerning scanner design and construction are available at orngm." stanford.edu/pbrown. Images were scanned at a ... resolution of 20 µm per pixel. A separate scan, using the appropriate excitation line, was done for each of the two fluorophores used. During the scanning process, the ratio between the signals in the two channets was calculated for several array elements containing total genomic DNA. To normalize the two channels with respect to overall intensity, we then adjusted photomultiplier and laser power settings such that the signal ratio at these elements was as close to 1.0 as possible. The combined images were analyzed with custom-written software. A bounding box, fitted to the size of the DNA spots in 'each quadrant, was placed over each array element. The average fluorescent intensity was calculated by summing the intensities of each pixel present in a bounding box, and then dividing by the total number of pixels. Local area background was calculated for each array element by determining the average fucrescent intensity for the lower 20% of pixel intensities. Although this method tends to underesting the background, causing an underestimation of extreme ratios, it produces a very consistent and noisetolerant approximation. Although the analog-todigital board used for data collection poss wide dynamic range (12 bits), several signals were saturated (greater than the maximum signal intensity allowed) at the chosen settings. Therefore, extrem ratios at bright elements are generally underestimated. A signal was deemed significant if the average intensity after background subtraction was at least 2.5-fold higher than the standard deviation in the background measurements for all elements on the
- 50. In addition to the 17 genes shown in Table 1, three additional genes were induced by an average of more than threefold in the duplicate experiments, but in one of the two experiments, the induction was less than twofold (range 1.6- to 1.9-fold)
- 51. We thank H. Bennett, P. Spellman, J. Ravetto, M. Eisen, R. Pillai, B. Dunn, T. Ferea, and other members of the Brown lab for their assistance and helpful advice. We also thank S. Friend, D. Botstein, S. Smith, J. Hudson, and D. Dolginow for advice, support, and encouragement; K. Struhl and S. Chatteree for the Tup1 deletion strain; L. Fernandes for helpful advice on Yap1; and S. Klapholz and the reviewers for many helpful comments on the manuscript. Supported by a grant from the National Hu-Genome Research Institute (NHGRI) (HG00450), and by the Howard Hughes Medical Institute (HHMI). J.D.R. was supported by the HHMI and the NHGRI. V.R. was supported in part by an Institutional Training Grant in Genome Science (T32 HG00044) from the NHGRI. P.O.B. is an associate investigator of the HHMI.

5 September 1997; accepted 22 September 1997

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Human Genome Placed on Chip; Biotech Rivals Put It Up for Sale

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The genome on a chip has arrived.

Melding high technology with biology, several companies are rushing to sell slivers of glass or nylon, some as small as postage stamps, packed with pieces of all 30,000 or so known human genes.

The new products will allow scientists to scan all genes in a human tissue sample at once, to determine which genes are active, a job that previously required two or more chips. The wholegenome chips will lower the cost and increase the speed of a widely used test that has transformed biomedical research in the last few years.

"It's sort of a milestone event, very similar to generating an integrated circuit of the genome," said Stephen P. A. Fodor, the chief executive of Affymetrix Inc., the leading seller of gene chips, which are also called microarrays.

Affymetrix, based in Santa Clara, Calif., is expected to announce today that it is accepting orders for its whole-genome chip.

The announcement seems timed to steal some thunder from the rival Agilent Technologies, which is based in nearby Palo Alto. Agilent is to be the host of an analyst meeting today and it plans to announce then that it has started shipping test versions of its whole-genome chip.

Applied Biosystems of Foster City, Calif., a unit of the Applera Corporation, started the race in July with an announcement that it would have a whole-genome chip out by the end of this year. NimbleGen Systems, a small company in Madison, Wis., announced a few days later that it had a genome on a chip that it was not selling but that it was using to run tests for customers.

Gene chips, which detect genes that are active, meaning they are being used to make a protein, have become essential tools. Scientists try to understand the genetic mechanisms of disease by seeing which genes are turned on in, say, a sick kidney or lung compared with those active in a healthy organ. Pharmaceutical companies look at gene activity patterns to try to predict the effects of drugs.

Scientists have found that tumors that look the same under the microscope can differ in terms of which genes are active. So studying gene patterns could become a way to discriminate between deadly and not-so-deadly tumors, or to predict which drug will work best for a particular patient.

Still, even some vendors conceded that the change from two chips to one is more symbolic than revolutionary.

"You can do just as good science with two chips, it costs you a little more," said Roland Green, the vice president for research and development at NimbleGen.

Some scientists questioned whether the chips really have all human genes, because the exact number and identities of all the genes is not known.

The advent of the genome on a chip is, however, evidence that biotechnology, to the extent that it uses electronics, is experiencing some of the rapid progress that has made semiconductors and computers continuously cheaper and smaller.

"One of the effects everyone is looking for in the genomics area is Moore's law -- more data, less money," said Doug Dolginow, an executive vice president at Gene Logic, which sells data from gene chip studies to pharmaceutical companies. "This is a step in that direction."

Moore's law states that the number of transistors on a semiconductor chip doubles every 18 months.

Affymetrix's gene chips are, in fact, made with the same techniques used to make semiconductor chips. In the mid-1990's, the company came out with a set of five chips covering what was then known of the human genome. After the human genome sequence was virtually completed in 2000, the company developed a two-chip set with all the known genes. Now it has the single chip, which some scientists say will be more convenient.

"We like to be able to look at all genes at one time to get a global view of what's going on," said John R. Walker, who runs gene chip operations at the Genomics Institute of the Novartis Research Foundation in San Diego.

Costs should also be lower. Gene chips have been so expensive that many academic scientists still make their own rather than buy them. Affymetrix said it would sell its whole-genome chips for \$300 to \$500 each, depending on volume, little more than half the price of the two-chip set. The other companies have not announced prices.

For Affymetrix, a successful whole-genome chip "is essential for them to maintain their dominance" of high-end microarrays, said Edward A. Tenthoff, an analyst at U.S. Bancorp Piper Jaffray. Affymetrix had total product sales in 2002 of about \$250 million, and a company spokesman said that human genome chips are its top-selling product.

Mr. Tenthoff, who recommends Affymetrix stock, said the company's sales growth rate had moderated as it faces tougher competition. Agilent, a spinoff of Hewlett-Packard that makes its gene chips by printing DNA components onto glass slides using ink jet printers, has gained share, he said. Applied Biosystems, the largest maker of genomics equipment over all, will be

entering the microarray segment of the business with its whole-genome chip, emphasizing the connection of that product to the others it offers, including the gene database developed by its sister company, Celera Genomics.

Jeffrey Trent, scientific director of the Translational Genomics Research Institute in Phoenix, said that while whole-genome chips are useful for medical discovery, the biggest growth of the market will be for chips that can be used by doctors to do diagnoses. And whole-genome chips are too cumbersome for that, he said. Rather, once scientists use the whole-genome chips to find particular genes that are associated with, say, tumor aggressiveness or drug effectiveness, he said, they will then make smaller and cheaper chips containing just those genes for use in diagnosis.

Agilent Technologies

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PALO ALTO, Calif., Oct. 2, 2003

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- **Archives**

Agilent Technologies Inc. (NYSE: A) today announced it has shipped whole human-genome microarrays to customers for testing and evaluation. The whole genome microarray is based on Agilent's new doubledensity format, which can accommodate 44,000 features on a single 1" x 3" glass-slide microarray. The new platform enables drug-discovery and disease researchers to perform whole-genome screening at a lower cost and with higher reproducibility.

"This is an important step toward our release of the first whole human-genome microarray product, which is expected to be available for order before the end of the year," said Barney Saunders, vice president and general manager of Agilent's BioResearch Solutions Unit. " Customers have long wanted a onesample, one-chip format with the increased sensitivity associated with 60-mer probes. The cost savings and high-quality performance make this product a compelling alternative for scientists who make their own microarrays."

Agilent's microarrays are based on the industry-standard 1" x 3" (25mm x 75mm) format, which is compatible with most commercial microarray scanners. All Agilent commercial microarrays are developed using content from public databases and proprietary sources, with full sequence and annotation information made available to customers. Gene sequences for probes are developed using algorithms and then validated empirically through iterative wet-lab testing procedures. The result is a microarray comprised of functionally validated probes, with the most up-to-date and comprehensive genome information commercially available.

Advantages of the double-density format include:

- Lower cost. Not only is one microarray less expensive than two, it requires fewer reagents and reduces instrumentation demands.
- Streamlined workflow. Researchers need prepare and process only one microarray instead of two. This also results in fewer steps in the subsequent data analysis.
- Greater reproducibility. Use of a single microarray further reduces unnecessary variability in experimental conditions.
- Smaller sample use. A smaller quantity of sample material is required to perform an experiment.

Availability

Agilent's Whole Human Genome Microarray is expected to be available for order by the end of the year.

About Agilent Technologies

Agilent Technologies Inc. (NYSE: A) is a global technology leader in communications, electronics, life sciences and chemical analysis. The company's 30,000 employees serve customers in more than 110 countries. Agilent had net revenue of \$6 billion in fiscal year 2002. Information about Agilent is available Search Agile

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Agilent | Agilent Technologies ships whole human genome on single microarray to gene e... Page 2 of 2

on the Web at www.agilent.com.

Forward-Looking Statements

This news release contains forward-looking statements (including, without limitation, statements relating to Agilent's expectation that its whole-genome microarray platform will be available for order before the end of 2003) that involve risks and uncertainties that could cause results to differ materially from management's current expectations. These and other risks are detailed in the company's filings with the Securities and Exchange Commission, including its Annual Report on Form 10-K for the year ended Oct. 31, 2002, its Quarterly Report on Form 10-Q for the quarter ended July 31, 2003 and its Current Report on Form 8-K filed Aug. 18, 2003. The company assumes no obligation to update the information in this press release.

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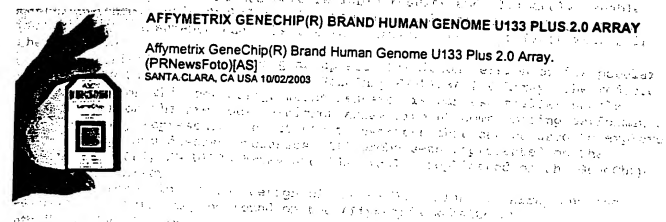
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Affymetrix GeneChip(R) Brand Human Genome U133 Plus 2.0 Array. (PRNewsFoto)[AS] The supersual transfer of the second of the procedure SANTA CLARA, CA USA 10/02/2003 one as to the exercise we can exist for an east the problem in each of

More Than 1 Million Probes Analyze Expression Levels of Nearly 50,000 RNA Transcripts and Variants on a Single Array the Size of a Thumbnail The second secon

SANTA CLARA, Calif., Oct. 2 /PRNewswire/ -- Affymetrix, Inc., (Nasdaq: AFFX) announced today that it is taking orders for its new GeneChip(R) brand Human Genome U133 Plus 2.0 Array, offering researchers the protein-coding content of the human genome on a single commercially available. catalog microarray. The HG-U133 Plus 2.0 Array analyzes the expression level of nearly 50,000 RNA transcripts and variants with 22 different probes per transcript, providing superior data quality unmatched by technologies using a single probe per transcript.

(Photo: http://www.newscom.com/cgi-bin/prnh/20031002/SFTH021)

"With about 1.3 million probes on a chip the size of a human thumbnail, the Human Plus Array represents a leap in array technology data capacity, and further demonstrates the unique power and potential of our technology to explore vast areas of the genome, " said Trevor J. Nicholls, Ph.D., Chief Commercial Officer. "Multiple independent measurements for each transcript ensure that our data quality remains the industry standard, even as our data capacity increases dramatically."

The HG-U133 Plus 2.0 Array, which will ship in October, combines the content of the previous HG-U133 two-array set with nearly 10,000 new probe sets representing about 6,500 new genes, for a total of nearly 50,000 RNA transcripts and variants. This new information, verified against the latest version of the publicly available genome map, provides researchers the most comprehensive and up-to-date genome-wide gene expression analysis. The probe design strategy of the HG-U133 Plus 2.0 Array is identical to the previous HG-U133 Set, providing very strong data concordance between the two products. With more than double the data capacity of the previous-generation Affymetrix human product, the HG-U133 Plus 2.0 Array can significantly cut processing and analysis time for scientists in the lab, freeing up valuable resources and accelerating research.

The HG-U133 Plus 2.0 Array sets a new standard for the number of genes and transcripts on any commercially available single array for human gene

expression analysis, while maintaining Affymetrix' unrivaled data quality. The HG-U133 Plus 2.0 Array uses 22 independent measures to detect the hybridization of each transcript on the array, 1.3 million data points in all, more than 30 times that of any other microarray technology. Using multiple, independent measurements provides optimal sensitivity and specificity, and the most accurate, consistent and statistically significant results possible. "More data points produce more reliable results and ultimately, enable better science," said Nicholls. "Our powerful probe set strategy gives our customers the assurance that their array results actually reflect what's in their sample."

Affymetrix is also launching an updated 11-micron version of its popular 18-micron HG-U133A Array called the GeneChip HG-U133A 2.0 Array. The reduced feature size on this new design means researchers can use smaller sample volumes than on the previous 18-micron array without compromising performance. This new array represents over 20,000 transcripts that can be used to explore human biology and disease processes. All probe sets represented on the original GeneChip HG-U133A Array are identically replicated on the GeneChip HG-U133A 2.0 Array.

More information on the design of the HG-U133 Plus 2.0 Array and the HG-U133A 2.0 Array may be found on the Affymetrix website at http://www.affymetrix.com.

Affymetrix will be presenting further information on this and other products at the BioTechnica trade show in Hanover, Germany on Oct. 7-9, 2003. The Company will also hold a press conference on Oct. 7, from 11 a.m. to 12 p.m. at the show regarding the new Human Genome U133 Plus 2.0 Array. If you would like to attend this press conference, please contact Caroline Stupnicka at c.stupnicka@northbankcommunications.com.

About Affymetrix:

Affymetrix is a pioneer in creating breakthrough tools that are driving the genomic revolution. By applying the principles of semiconductor technology to the life sciences, Affymetrix develops and commercializes systems that enable scientists to improve the quality of life. The Company's customers include pharmaceutical, biotechnology, agrichemical, diagnostics and consumer products companies as well as academic, government and other non-profit research institutes. Affymetrix offers an expanding portfolio of integrated products and services, including its integrated GeneChip platform, to address growing markets focused on understanding the relationship between genes and human health. Additional information on Affymetrix can be found at http://www.affymetrix.com.

All statements in this press release that are not historical are "forward-looking statements" within the meaning of Section 21E of the Securities Exchange Act as amended, including statements regarding Affymetrix' "expectations," "beliefs," "hopes," "intentions," "strategies" or the like. Such statements are subject to risks and uncertainties that could cause actual results to differ materially for Affymetrix from those projected, including, but not limited to risks of the Company's ability to achieve and sustain higher levels of revenue, higher gross margins, reduced operating expenses, uncertainties relating to technological approaches, manufacturing, product development, market acceptance (including uncertainties relating to product development and market acceptance of the GeneChip HG-Ul33 Human Plus 2.0 Array and the HG-U133A 2.0), personnel retention, uncertainties related to cost and pricing of Affymetrix products, dependence on collaborative partners, uncertainties relating to sole source suppliers, uncertainties relating to FDA and other regulatory approvals, competition, risks relating to intellectual property of others and the uncertainties of patent protection and litigation. These and other risk factors are discussed in Affymetrix' Form 10-K for the

year ended December 31, 2002 and other SEC reports, including its Quarterly Reports on Form 10-Q for subsequent quarterly periods. Affymetrix expressly disclaims any obligation or undertaking to release publicly any updates or revisions to any forward-looking statements contained herein to reflect any change in Affymetrix' expectations with regard thereto or any change in events, conditions, or circumstances on which any such statements are based.

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Macroresults through Microarrays

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The third enactment of Cambridge Healthtech Institute's Macroresults through Microarrays meeting was held in Boston (MA, USA) from 29 April—1 May 2002. The subtheme of this year's meeting was 'advancing drug discovery', a widely touted application for array technology.

The evolution of microarrays

If you were asked 'Who first conceived.' of the idea of microarrays', who would come to mind? Mark Schena perhaps, first author of the seminal 1995 paper on cDNA arrays [1]? Maybe Pat Brown, Schena's then supervisor? Or perhaps Stephen Fodor, the primary driver behind Affymetrix's (http://www. affymetrix.com) oligonucleotide-based platform [2]. Brits might even chant the name of Ed Southern [3]. Well, according to Roger Ekins (University College London Medical School; http://www. ucl.ac.uk/medicine/) all these answers would be wrong. It was in fact Ekins and his colleagues who first conceived of and patented 'a new generation of ultrasensitive, miniaturized assays for protein and DNA-RNA measurement based on the use of microarrays' in the mid 1980s [4]. The concept and potential of array technology was more fully described in a later publication, in which Ekins et al. [5] concluded that antibody microspots of -50 µm² could be achieved, and that as many as 2 million different immunoassays could, in principle, be accommodated on a surface area of 1 cm².

Technological innovation In practice, it took a different biological molecule (DNA), a different research

group, and a leap into microfabrication technology to even begin approaching these kinds of densities -[Affymetrix patent 6045996 talks of one million spots cm-2]. Of course, advancing technology is one of the driving engines behind the genomics juggernaut, and we are already seeing '4th generation' machines for fabricating DNA chips. If the company representatives at this meeting are to be believed (and their cases seemed strong), spotting is out, and in situ fabrication of oligonucleotide-based 'iterative custom arrays' is in. Whether you go with the Combimatrix's (http:// www.combimatrix.com) electrochemically directed synthesis and detection system, febit's (http://www.febit.com) Geniom® technology, or Nimblegen's (http://www.nimblegen.com) Maskless Array Synthesizer technology is a matter of personal choice. However, each of these machines provides the flexibility to design variable length oligonucleotide probes from sequences inputted by the user, and then perform in situ synthesis of an array. Each system also boasts unique advantages. For example, Combimatrix's biological array processor is a semiconductor coated with a 3D layer of porous material in which DNA, RNA, peptides or small molecules can be synthesized or immobilized within discrete test sites, while febit's Geniom One® is a fully integrated gene-expression analysis system with minimal user hands-on time - the probe sequences are programmed, the RNA samples inserted, and the gene expression data is pumped out a few hours later.

Cell- and tissue-based arrays

Array technology is in most people's: minds firmly linked with gene-expression profiling. Fewer are aware that cell- and tissue-based arrays have been developed, and how they can provide a vital extra dimension to research. In support of this, Barry Bochner gave an update on the cell-based array system that Biolog (http://www.biolog.com) has produced for simultaneously measuring the effects of one gene in the cell under thousands of growth conditions 1... (see [6] for further details). David Walt (Tufts University; http://www.tufts. edu/) is developing single live cell ar- :-rays using optical imaging fiber (OIF) technology. An array of microwells is fabricated on the face of an OIF at densities of up to 10 million wells cm-2. Cells are then added to the wells and disperse at an average of one cell per well. Physiological and genetic responses of each cell are measured via fluorescence produced by reporter genes (e.g. locZ, gfp. Assays performed so far include yeast live or dead cell assay, microenvironment pH and O₂ measurements, promotor responses using the lacZ and phoA reporter genes, and protein-protein interactions using the yeast two-hybrid system. The main advantage of this system is that the cells remain alive during the assay, which means a real-time timecourse can be performed and/or the array passed from sample to sample. This would be useful in, for example, the scanning of a combinatorial drug library for specific physiological effects.

Tissue arrays are a useful complementary technology to DNA arrays because they can be used to help validate and

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understand the biological and medical significance of gene changes discovered using standard DNA arrays. For example, an array of tumor tissues can be screened for the protein (using immunohistochemistry), message (using in situ hybridization) and copy number (using comparative genomic hybridization) of a gene of interest, to determine if expression of the gene (or lack thereof) is related in any way to survival. They can also be used to predict the probability of clinical failure of lead compounds as a result of toxicity by evaluating the distribution of the drugtargets in normal tissue. Spyro Mousses and his co-workers at the National Human Genome Research Institute (http://www.nhgri.nih.gov/index.html) have built such arrays, including a multi-tumor array (-5000 specimens, and sections from 36 normal and 800 metastatic tissues) and a normal tissue array (76 tissue and 332 cell types).

The problem with proteins

It has been said that genomics tells us what might happen, transcriptomics indicates what should happen, and proteomics shows what is happening. The impact of functional proteomics on pharmaceutical R&D is rapidly increasing, and protein arrays are being used increasingly in both basic and applied research. Their use lies not only in comparative protein expression and interaction profiling, but also in diagnostics and drug discovery. However, an increasing number of researchers have found that protein arrays, like their cousins the DNA arrays, present several practical obstacles relating to their production and use. For example, in using Escherichia coli to produce recombinant eukaryotic proteins from a single expression vector, multiple protein products are often produced, suggesting mixes of truncated or otherwise altered proteins. There is also the obvious concern that the proteins might not be modified in a similar manner to

method for depositing and binding proteins to the selected substrate is yet to be determined, as is the best way to ensure that they are bound in a correctly folded, active conformation.

Several companies have been addressing these problems. Prolinx (http:// www.prolinxinc.com) is one such company, and Karin Hughes described their Versalinx™ chemistry for producing protein, peptide and small-molecule arrays. VersalinxTM uses solution-phase conjugation followed by immobilization, resulting in functional orientation of proteins and peptides on the substrate surface. It also offers the valuable additional benefit of exhibiting low non-specific binding. Sense Proteomic (http://www.senseproteomic.com) is also among those addressing these problems to develop robust protein arrays for drug discovery and clinical applications and has developed functional protein array formats based on specific disease tissues. Subtractive hybridization is used to identify genes with altered expression in breast tumor and cystic fibrosis compared to normal tissue. A high throughput cloning strategy (COVET™) is then used to produce libraries of genes that are tagged, cloned, expressed, purified and finally immobilized on glass slides. Initial validation studies have shown that the vast majority of the immobilized proteins do indeed retain biological function.

Stefan Schmidt and his company (GPC Biotech; http://www.gpcbiotech. de) have moved past the platform development stage and, with their focus firmly on drug discovery, are currently developing kinase-profiling arrays. Kinases are important targets for pharmaceutical drug discovery and therapy, and GPC's aim is to simultaneously detect multiple kinases, obtain activity profiles for different cell types, or analyze the ability of drug candidates to inhibit kinase activity. To do this, recombinant kinase substrates are immobilized on

mal rmembranes, incubated with purified ing kinase, and the substrates measured for the degree of phosphorylation.

Summary

Meetings like this, packed with exciting discoveries and intriguing and interesting innovation, heavily emphasize the pace at which biotechnology is advancing, to the extent that the number of options for genomic and proteomic researchers can become overwhelming. Although data analysis is perhaps the greatest current concern for array users, an increasing challenge will be to determine the approaches and technology that really work, and to do it in a timely manner.

References

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 Schena, M. et al. (1995) Quantitative monitoring of gene expression patterns with a complementary DNA microarray. Science 270, 467-470

11 4 6

- 2 Fodor, S.P. et al. (1991) Light-directed, spatially addressable parallel chemical synthesis. Science 251, 767-773
- 3 Southern, E.M. et al. (1992) Analyzing and comparing nucleic acid sequences by hybridization to arrays of oligonucleotides: evaluation using experimental models. Genomics 13, 1008–1017
- 4 Ekins, R.P. (1987) US Patent Application 8 803 000
- 5 Ekins, R. et al. (1989) High specific activity chemiluminescent and fluorescent markers: their potential application to high sensitivity and 'multi-analyte' immunoassays. J. Biolum. Chemilum. 4, 59-78
- 6 Rockett, J.C. (2002) Chip, chip, array! Three chips for post-genomic research. *Drug Discov. Today* 7, 458–459

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A two-dimensional gel database of rat liver proteins Jean-Paul Holmann useful in gene regulation and drug effects studies Norman G. Anderson A see the see present of seed and the seed of the company of the groups of the seed of th

Large Scale Biology Corporation.

A standard two-dimensional (2-D) protein map of Fischer 344 rat liver Rockville, MD

(F344MST3) is presented, with a tabular listing of more than 1200 protein species. Rockville, MD Sodium dodecy sulfate (SDS) molecular mass and isoelectric point have been established, based on positions of numerous internal standards. This map has been Lal et al., 09/002;485, filed December 31, 1997

(PF-0459)

used to connect and compare hundreds of 2-D gels of rat liver samples from a variety of studies, and forms the nucleus of an expanding database describing rat liver proteins and their regulation by various drugs and toxic agents. An example Exhibit "L" attached to Declaration of John C. of such's study, involving regulation of cholesterol synthesis by cholesterol-lower-Exhibit "L" attached to Declaration of Solar C.

Of such a study, involving regulation of cholesterol synthesis by cholesterol-lowerRockett, Ph.D.:

Ing drugs and a high-cholesterol diet, is presented. Since the map has been obtained with a widely used and highly reproducible 2-D gel system (the Iso-Dalt*) tained with a widely used and highly reproducible 2-D gel system (the Iso-Daltesystem), it can be directly related to an expanding body of work in other laboratories. The first of the second of the

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Abbrevistions: CBB, Coomassie Brilliant Blue; CPK, creatine phosphokinase; 2-D, two-dimensional; IEF, isoelectric focusing; MSN, master spot number; NP-40, Nonidet P-40, SDS, sodium dodecyl sulfate

1. Introduction in an argument of the fact of the contract of

and the first of a stationing carefus High-resolution two-dimensional electrophoresis of proteins, introduced in 1975 by O'Farrell and others [1-4], has been used over the ensuing 16 years to examine a wide variety of biological systems, the results appearing in more than 5000 published papers. With the advent of computerized systems for analyzing two-dimensional (2-D) gel images and constructing spot databases, it is also possible to plan and assemble integrated bodies of information describing the appearance and regulation of thousands of protein gene products [5, 6]. Creating such databases involves amassing and organizing quantitative data from thousands of 2-D gels, and requires a substantial commitment in technology and resources.

Given the long-term effort required to develop a protein database, the choice of a biological system takes on considerable importance. While in vitro systems are ideal for answering many experimental questions, especially in cancer research and genetics, our experience with cell cultures and tissue samples suggests that some in vivo approaches could have major advantages. In particular, we have noticed that liver tissue samples from rats and mice appear to show greater quantitative reproducibility (in terms of individual protein expression) than replicate cell cultures. This is perhaps a natural result of the homeostasis maintained in a complete animal vs. the well-known variability of cell cultures, the latter due principally to differences in reagents (e.g., fetal bovine serum), conditions (e.g., pH) and genetic "evolution" of cell lines while in culture. It is also more difficult to generate adequate amounts of protein from cell culture systems (particularly with attached cells), forcing the invesligator to resort to radioisotope-based or silver-based staindetection methods. While these methods are more sensitive (sometimes much more sensitive) than the Coomassie Brilliant Blue (CBB) stain typically used for protein detection in "large" protein samples, they are generally more variable, more labor-intensive and, in the case of radiographic methods, may generate highly "noisy" images, due to the properties of the films used. By contrast, large protein samples can easily be prepared from liver using urea/Nonidet P-40 (NP-40) solubilization and stained with CBB, which has the advantage of being easily reproducible [8]. Finally, there remains the question of the "truthfulness" of many in vitro systems as compared to their in vivo analogs; how great are the changes caused by the introduction into a culgroupe with Eggreg to 18 ft at 122

ture and the associated shift to strong selection for growth, and how do these affect experimental outcomes? Hence the apparent advantages of in vitro systems, in terms of experimental manipulation, may be counterbalanced by other factors relating to 2-D data quality.

Source State State of the State of the There is a second important class of reasons for exploring the use of an in vivo biological system such as the liver. Historically, there have been two broad approaches to the mechanistic dissection of biochemical processes in intact cellular systems: genetics (a search for informative mutants) and the use of chemical agents (drugs and chemical toxins). Both approaches help us to understand complex systems by disrupting some specific functional element and showing us the result. With the development of techniques for genetic manipulation and cloning, the genetic approach can be effectively applied either in vitro or in vivo, although the in vitro route is usually quicker. The chemical approach can also be applied to either sort of biological system; here, however, the bulk of consistently acquired information is in experimental animals (rats and mice). While most biologists know a short list of compounds having specific, experimentally useful effects (e.g., inhibitors of protein synthesis, ionophores, polymerase inhibitors, channel blockers, nucleotide analogs, and compounds affecting polymerization of cytoskeletal proteins), there is a much larger number of interesting chemically-induced effects, most of them characterized by toxicologists and pharmacologists in rodent systems. Just as a thorough genetic analysis would involve saturating a genome with mutations, it is possible to imagine a saturating number of drugs, the analysis of whose actions would reveal the complete biochemistry of the cell. While organized drug discovery efforts usually target specific desired effects, the nature of the process, with its dependence on screening large numbers of compounds, necessarily produces many unanticipated effects. It is therefore reasonable to suppose that the required broad range of compounds necessary to achieve "biochemical saturation" may be forthcoming; in fact, it may already exist among the hundreds of thousands of compounds that failed to qualify as drugs.

Among organs, the liver is an obvious choice for the study of chemical effects because of its well-known plasticity and responsiveness. The brain appears to be quite plastic (e.g. [7]), but it is a complicated mixture of cell types requiring skillful dissection for most experiments. The kidney, while quite responsive, also presents a potentially confounding mixture of cell types. The liver, by contrast, is made up of one predominant cell type which is easy to solubilize: the hepatocyte, representing more than 95% of its mass. Most importantly, the liver performs many homeostatic functions that require rapid modulation of gene expression. It appears that most chemical agents tested affect gene expression in the liver at some dosage (N. Leigh Anderson, unpublished observations), an interesting contrast to our earlier work with lymphocytes, for example, which seem to be much less responsive. Such results conform to the expectation that cells with a homeostatic, physiological role should be more plastic than cells differentiated for a purpose dependent on the action of a limited number of specific genes.

The liver also allows the parallels between in vitro and in vivo systems to be examined in detail. Significant progress

has been made in the development of mouse, rat and human hepatocyte culture systems, as well as in precision-cut tissue slices. Using such an array of techniques, it is possible to assemble a/matrix of mammalian systems including mouse and rat in vivo on one level and mouse, rat and human high on a second level, and to compare effects between species and between systems. This approach allows us to draw informed conclusions regarding the biochemical universality of biological responses among the mammals, and to offer some insight into the validity of in vitro approaches for toxicological screening. We believe this data will be necessary if in vitro alternatives are to achieve wide usage in government-mandated safety testing of drugs, consumer products and industrial and agricultural chemicals.

A number of interesting studies have been published using 2-D mapping to examine effects in the rodent liver. A number of investigarors have made use of the technique to screen for existing genetic variants [8-11] or induced mutations [12-14], mainly in the mouse. This work builds on the wealth of genetic information available on the mouse and its established position as a mammalian mutation-detection system. While some studies of chemical effects have been undertaken in the mouse [15-17], most have used the rat [18-23]. The examination of the cytochrome p-450 system, in particular, has been carried out almost exclusively on the rat [24, 25].

These considerations lead us to conclude that rodent liver offers the best opportunity to systematically examine an array of gene regulation systems, and ultimately to build a predictive model of large-scale mammalian gene control. The basic underlying foundation of such a project is a reliable, reproducible master 2-D pattern of liver, to which ongoing experimental results can be referred. In this paper, we report such a master pattern for the acidic and neutral proteins of rat liver (pattern F344MST3). In future, this master will be supplemented by maps of basic proteins, and analogous maps of mouse and human liver.

2 Materials and methods

2.1 Sample preparation

Liver is an ideal sample material for most biochemical studies, including 2-D analysis. A sample is taken of approximately 0.5 g of tissue from the apical end of the left lobe of the liver. Solubilization is effected as rapidly as practical; a delay of 5-15 min appears to cause no major alteration in liver protein composition if the liver pieces are kept cold (e.g., on ice) in the interim. In the solubilization process, the liver sample is weighed, placed in a glass homogenizer (e.g., 15 mL Wheaton); 8 volumes of solubilizing solution.

The solubilizing solution is composed of 2% NP-40 (Sigma), 9 m urea (analytical grade, e.g., BDH or Bio-Rad), 0.5% dithiothreitol (DTT; Sigma) and 2% carrier ampholytes (pH 9-11 LKB: these come as a 20% stock solution, so 2% final concentration is achieved by making the final solution 10% 9-11 Ampholine by volume). A large batch of solubilizer (several hundred mL) is made and stored frozen at -80°C in aliquots sufficient to provide enough for one day's estimated sample preparation requirement. The solution is never allowed to become warmer than room temperature at any stage during preparation or thawing for use, since heating of concentrated urea solutions can produce contaminants that covalently modify proteins producing artifactual charge shifts. Once thawed, any unused solubilizer is discarded.

is added (i.e., 4 mL per 0.5 g tissue) and the mixture is homogenized using first the loose- and then then the tight-fitting glass pestle. This takes approximately 5 strokes with each pestle and is carried out at room temperature because urea would crystallize out in the cold. Once the liver sample is thoroughly homogenized in the solubilizer, it is assumed that all the proteins are denatured (by the chaotropic effect of the urea and NP-40 detergent) and the enzymes inactivated by the high pH (-9.5). Therefore these samples may be kept at room temperature until they can be centrifuged or frozen as a group (within several hours of preparation). The samples are centrifuged for 6 × 10⁴ g min (e.g., 500 000 × g for 12 min using a Beckman TL-100 centrifuge). The centrifuge rotor is maintained at just below room temperature (e.g., 15-20°C), but not too cold, so as to prevent the precipitation of urea. The centrifuge of choice is a Beckman TL-100 because of the sample tube sizes available, but any ultracentrifuge accepting smallish tubes will suffice. When an appropriate centrifuge is not available near the site of sample preparation, samples can be frozen at -80°C and thawed prior to centrifugation and collection of supernatants. Each supernatant is carefully removed following centrifugation and aliquoted into at least 4 clean tubes for storage. This is done by transferring all the supernatant to one clean tube, mixing this gently (to assure homogeneous composition) and then dividing it into 4 aliquots. The aliquots are frozen immediately at -80°C. These multiple aliquots can provide insurance against a failed run or a freezer breakdown.

2.2 Two-dimensional electrophoresis

Sample proteins are resolved by 2-D electrophoresis using the 20 × 25 cm Iso-Dait[®] 2-D gel system ([26-29]; produced by LSB and by Hoefer Scientific Instruments, San Francisco) operating with 20 gels per batch. All first-dimensional isoelectric focusing (IEF) gels are prepared using the same single standardized batch of carrier ampholytes (BDH 4-8A in the present case, selected by LSB's batchtesting program for rat and mouse database work**). A 10 ul sample of solubilized liver protein is applied to each gel, and the gels are run for 33 000 to 34 500 volt-hours using a progressively increasing voltage protocol implemented by a programmable high-voltage power supply. An Angelique" computer-controlled gradient-casting system (produced by LSB) is used to prepare second-dimensional sodium dodecyl sulfate (SDS) polyacrylamide gradient slab gels in which the top 5% of the gel is 11%T acrylamide, and the lower 95% of the gel varies linearly from 11% to 18%T.

This system has recently been modified so as to employ a commercially available 30.8%T acrylamide/N,N-methylenebisacrylamide prepared solution (thus avoiding the handling of the solid acrylamide monomer) and three additional stock solutions: buffer (made from Sigma pre-set Tris), persulfate and N,N,N,N-tetramethylethylenediamine (TEMED). Each gel is identified by a computer-printed filter paper label polymerized into the lower left corner of the gel. First-dimensional IEF tube gels are loaded

directly (as extruded) onto the slab gels without equilibration, and held in place by polyester fabric wedges (Wedgies", produced by LSB) to avoid the use of hot agarose. Second-dimensional slab gels are run overnight, in groups of 20, in cooled DALT tanks (10°C) with buffer circulation. All run parameters, reagent source and lot information, and notations of deviation from expected results are entered by the technician responsible on a detailed, multi-page record of the experiment.

2.3 Staining

Following SDS-electrophoresis, slab gels are stained for protein using a colloidal Coomassie Blue G-250 procedure in covered plastic boxes, with 10 gels (totalling approximately 1 L of gel) per box. This procedure (based on the work of Neuhoff [30, 31]) involves fixation in 1.5 L of 50% ethanol and 2% phosphoric acid for 2h, three 30 min washes, each in 2L of cold tap water, and transfer to 1-5L of 34% methanol, 17% ammonium sulfate and 2% phosphoric acid for 1 h, followed by the addition of a gram of powdered Coomassie Blue G-250 stain. Staining requires approximately 4 days to reach equilibrium intensity, whereupon gels are transferred to cool tap water and their surfaces rinsed to remove any particulate stain prior to scanning. Gels may be kept for several months in water with added sodium azide. The water washes remove ethanol that would dissolve the stain (and render the system noncolloidal, with high backgrounds). The concentrated ammonium sulfate and methanol solution is diluted by equilibration with the water volume of the gels to automatically achieve the correct final concentrations for colloidal staining. Practical advantages of this staining approach can be summarized as follows: (i) the low, flat background makes computer evaluation of small spots (max OD < 0.02) possible, especially when using laser densitometry; (ii) up to 1500 spots can be reliably detected on many gels (e.g., rat liver) at loadings low enough to preserve excellent resolution; and (iii) reproducibility appears to be very good: at least several hundred spots have coefficients of reproducibility less than 15%. This value is at least as good as previous CBB methods, and significantly better than many silver stain systems.

2.4 Positional standardization

The carbamylated rabbit muscle creatine phosphokinase (CPK) standards [32] are purchased from Pharmacia and BDH. Amino acid compositions, and numbers of residues present in proteins used for internal standardization, are taken from the Protein Identification Resource (PIR) sequence database [33].

2.5 Computer analysis

Stained slab gels are digitized in red light at 134 micron resolution, using either a Molecular Dynamics laser scanner (with pixel sampling) or an Eikonix 78/99 CCD scanner. Raw digitized gel images are archived on high-density DAT tape (or equivalent storage media) and a greyscale videoprint prepared from the raw digital image as hard-copy backup of the gel image. Gels are processed using the Kepler software system (produced by LSB), a commercially available workstation-based software package built on

^{**} This material (succeeding certified batches of which are available from Hoefer Scientific Instruments) has the most linear pH gradient produced by any ampholyte tested except for the Pharmacia wide range (which has an unacceptable tendency to bind high-molecular weight acidic proteins, causing them to streak).

some of the principles of the earlier TYCHO system [34-41]. Procedure PROC008 is used to yield a spotlist giving position, shape and density information for each detected spot. This procedure makes use of digital filtering, mathematical morphology techniques and digital masking to remove the background, and uses full 2-D least-squares optimization to refine the parameters of a 2-D Gaussian shape for each spot. Processing parameters and file locations are stored in a relational database, while various log files detailing operation of the automatic analysis software are archived with the reduced data. The computed resolution and level of Gaussian convergence of each gel are inspected and archived for quality control purposes.

Experiment packages are constructed using the Kepler experiment definition database to assemble groups of 2-D patterns corresponding to the experimental groups (e.g., treated and control animals). Each 2-D pattern is matched to the appropriate "master" 2-D (pattern (pattern F344MST3 in the case of Fischer 344 rat liver), thereby providing linkage to the existing rodent protein 2-D databases. The software allows experiments containing hundreds of gels to be constructed and analyzed as a unit, with up to 100 gels displayed on the screen at one time for comparative purposes and multiple pages to accommodate experiments of > 1000 gels. For each treatment, proteins showing significant quantitative differences vs. appropriate controls are selected using group-wise statistical parameters (e.g., Student's t-test, Kepler procedure STUDENT). Proteins satisfying various quantitative criteria (such as P< 0.001 difference from appropriate controls) are represented as highlighted spots onscreen or on computer-plotted protein maps and stored as spot populations (i.e., logical vectors) in a liver protein database. Quantitative data (spot parameters, statistical or other computed values) are stored as real-valued vectors in the database. Analysis of coregulation is performed using a Pierson product-moment correlation (Kepler procedure CORREL) to determine whether groups of proteins are coordinately regulated by any of the treatments. Such groups can be presented graphically on a protein map, and reported together with the statistical criteria used to assess the level of coregulation. Multivariate statistical analysis (e.g., principal components' analysis) is performed on data exported to SAS (SAS Institute).

2.6 Graphical data output

Graphical results are prepared in GKS and translated within Kepler® into output for any of a variety of devices. Linedrawing output is typically prepared as Postscript and printed on an Apple Laserwriter. Detailed maps presented here have been generated using an ultra-high-resolution Postscript-compatible Linotronic output device. Greyscale graphics are reproduced from the workstation screen using a Seikosha videoprinter. Patterns are shown in the standard orientation, with high molecular mass at the top and acidic proteins to the left.

2.7 Experiment LSBC04

In the study described here 12-week-old Charles River male F344 rats were used. Diets were prepared at LSB, based on a Purina 5755M Basal Purified Diet. Lovastatin and cholestyramine were obtained as prescription pharma-

ceuticals, ground and mixed with the diet at concentrations of 0.075% and 1%, respectively. The high cholesterol diet was Purina 5801M-A (5% cholesterol plus 1% sodium cholate in the control diet). Animal work was carried out by Microbiological Associates (Bethesda, MD). Animals were acclimatized for one week on the control diet, fed test or control diets for one week, and sacrificed on day 8. Average daily doses of lovastatin and cholestyramine in appropriate groups were 37 mg/kg/day and 5 g/kg/day, respectively, based on the weight of the food consumed. Liver samples were collected and prepared for 2-D electrophores is according to the standard liver protocol (homogenization in 8 volumes of 9 m urea, 2% NP-40, 0.5% dithiothreitol, 2% LKB pH-9-11 carrier ampholytes, followed by centrifugation for 30 min at 80 000 × g). Kidney, brain and plasma samples were frozen. Gels were run as described above, and the data was analyzed using the Keplers, system. Gels were scaled, to remove the effect of differences in protein loading, by setting the summed abundances of a large number of matched spots equal for each gel (linear scaling).

3 Results and discussion

3.1 The rat liver protein 2-D map

F344MST3 is a standard 2-D pattern of rat liver proteins, based on the Fischer 344 strain. This pattern was initiated from a single 2-D gel and extensively edited in an experiment comparing it to a range of protein loads, so as to include both small spots and well-resolved representations of high-abundance spots. More than 700 rat liver 2-D patterns have been matched to F344MST3 in a series of drug effects and protein characterization experiments, and numerous new spots (induced by specific drugs, for instance) have been added as a result. A modified version including additional spots present in the Sprague-Dawley outbred rat has also been developed (data not shown). Figure 1 shows a greyscale representation and Fig. 2 a schematic plot of the master pattern. More than 1200 spots are included, most of which are visible on typical gels loaded with 10 µL of solubilized liver protein prepared by the standard method and stained with colloidal Coomassie Blue. Master spot numbers (MSN's) have been assigned to all proteins, and appear in the following figures, each showing one quadrant of the pattern. Figure 3 shows the upper left (acidic, high molecular mass) quadrant, Fig. 4 the upper right (basic, high molecular mass) quadrant, Fig. 5 the lower left (acidic, low molecular mass) quadrant, and Fig. 6 the lower right (basic, low molecular mass) quadrant. The quadrants overlap as an aid to moving between them. The gel position (in 100 micron units), isoelectric point (relative to the CPK internal pl standards) and SDS molecular mass (from the calibration curve in Fig. 8) are listed for each spot (Table 1). Because of the precision of the CPK-pl values, these parameters can be used to relate spot locations between gel systems more reliably than using pl measurements expressed as pH. A major objective of current studies is the identification of all major spots corresponding to known liver proteins, as well as rigorous definitions of subcellular organelle contents. Of particular interest to us is the parallel development of identifications in the rat and mouse liver maps, allowing detailed comparisons of gene expression effects in the two systems. The results of these studies will be presented systematically in a later edition of this database,

but we include here a useful series of 22 orienting identifications as an aid to other users of the rat liver pattern (Table 2).

3.2 Carbamylated charge standards, computed p/s and molecular mass standardization

We have previously shown that the use of a system of closely-spaced internal pI markers (made by carbamylating a basic protein) offers an accurate and workable solution to the problem of assigning positions in the pI dimension [32]. The same system, based on 36 protein species made by carbamylating rabbit muscle CPK, has been used here to assign pI's to most rat liver acidic and neutral proteins. The standards were coelectrophoresed with total liver proteins, and the standard spots added to a special version of the master pattern F544MST3. The gel X-coordinates of all liver protein spots lying within the CPK charge train were then transformed into CPK pI positions by interpolation between the positions of immediately adjacent standards (Table 1) using a Kepler³ vector procedure.

It has proven possible to compute fairly accurate pl values for many proteins from the amino acid composition [42]. We have attempted here to test a further elaboration of this approach, in which we computed pf's for the CPK standards themselves, based on our knowledge of the rabbit muscle CPK sequence and the fact that adjacent members of the charge train typically differ by blockage of one additional lysine residue (Table 3). We compared these values to similar computed pl's for an additional set of carbamylated standards made from human hemoglobin beta chains and a series of rat liver and human plasma proteins of known position and sequence (Fig. 7, Table 4). The result demonstrates good concordance between these systems. Two proteins show significant deviations: liver fatty-acid binding protein (FABP; #1 in Table 4) and protein disulphide isomerase (#20 in the table). The FABP spot present on F344MST3 may represent a charge-modified version of a more basic parent spot closer to the expected pl, not resolved in the IEF/SDS gel. Of particular importance is the fact that, by comparing computed pros of sequenced but unlocated proteins with the CPK p/s, we can assign a probable gel location without making any assumptions regarding the actual gel pH gradient. This offers a useful shortcut, given the vagaries of pH measurement on small diameter IEF gels. We have used this approach to compute the CPK prs of all rat and mouse proteins in the PIR sequence database, as an aid to protein identification (data not shown).

In order to standardize SDS molecular weight (SDS-MW), we have used a standard curve fitted to a series of identified proteins (Fig. 8). Rather than using molecular mass per se, we have elected to use the number of amino acids in the polypeptide chain, as perhaps a better indication of the length of the SDS-coated rod that is sieved by the second dimension slab. The resulting values were multiplied by 112 (the weighted average mass of amino acids in sequenced proteins) to give predicted molecular masses. Because we use gradient slabs, we have not constrained the fitted curve to conform to any predetermined model; rather we tried many equations and selected the best using the program "Tablecurve" on a PC. The equation chosen was y = a + bx + c/x, where y is the number of residues, x is the gel

Y coordinate, a is 511.83, b is -0.2731 and c is 33183801. The resulting fit appears to be fairly good over a broad range of molecular mass.

3.3 An example of rat liver gene regulation: Cholesterol metabolism

Experiment LSBC04 was designed as a small-scale test of the regulation of cholesterol metabolism in vivo by three agents included in the diet: lovastatin (Mevacora, an inhibitor of HMG-CoA reductase); cholestyramine (a bile acid sequestrant that has the effect of removing cholesterol from the gut-liver recirculation); and cholesterol itself. The first two agents should lower available cholesterol and the third should raise it, allowing manipulation of relevant gene expression control systems in both directions. Such an experiment offers an interesting test of the 2-D mapping, system since most of the pathway enzymes are present in low abundance, many are membrane-bound and difficult, to solubilize, and the pathway itself is complex. Approximately 1000 proteins were separated and detected in liver homogenates. Twenty-one proteins were found to be affected by at least one treatment, and these could be divided intoseveral coregulated groups.

3.3.1 MSN 413 (putative cytosolic HMG-CoA synthase) and sets of spots regulated coordinately or inversely

State Same One group of spots (including a spot assigned to the cytosolic HMG-CoA synthase, MSN 413) showed the expected increase in abundance with lovastatin or cholestyramine, the synergistic further increase with lovastatin and cholestyramine, and a dramatic decrease with the high cholesterol diet. Spot number 413 is the most strongly regulated protein in the present experiment, showing a 5- to 10-fold induction after a 1 week treatment with 0.075 % lovastatin and 1% cholestyramine in the diet (Figs. 9 and 10). Its expression follows precisely the expectation for an enzyme whose abundance is controlled by the cholesterol level; it is progressively increased from the control levels by cholestyramine, lovastatin and lovastatin plus cholestyramine, and it sinks below the threshold of detection in animals fed the high cholesterol diet. This spot has been tentatively identified as the cytosolic HMG-CoA synthase, based on a reaction with an antiserum to that protein provided by Dr. Michael Greenspan at Merck Sharp & Dohme Research Laboratories. This enzyme lies immediately before HMG-CoA reductase in the liver cholesterol biosynthesis pathway, and is known to be co-regulated with it. Spot 413 has an SDS molecular weight of about 54 000 and a CPK pl of -11.4, in reasonably close agreement with a molecular weight of 57300 and a CPK pl of -15.7 computed from the known sequence of the hamster enzyme [43].

Using a classical product-moment correlation test (Kepler procedure CORREL), a series of five additional spots was found to be coregulated with 413. The level of correlation was exceedingly high (> 95%). Two of these, 1250 and 933, are at similar molecular weights and approximately one charge more acidic than 413 (Fig. 9), indicating that they may be covalently modified forms of the 413 polypeptide. This suspicion is strengthened by the observation that both spots are also stained by the antibody to cytosolic HMG-CoA synthase. The remaining three correlated spots appear

to comprise an additional related pair (1253 and 1001) of around 40 kDa and a single spot (1119) of around 28 kDa. Because these two presumed proteins are present at substantially lower abundances than 413, and because the cytosolic HMG-CoA synthase is reported to consist of only one type of polypeptide, they are likely to represent other, very tightly coregulated enzymes. A second group of six spots was selected based on a regulatory pattern close to the inverse of that for spot 413 (MSN's 34, 79, 178, 182, 204, 347; data not shown). For these proteins, the lowest level of expression occurs with exposure to lovastatin plus cholestyramine and the highest level upon exposure to the high-cholesterol diet. Spots 182 and 79 are highly correlated and lie about one charge apart at the same molecular weight; they may thus be isoforms of a single protein. The other four spots probably represent additional enzymes or subunits.

3.3.2 MSN 235 and coregulated spots

A third group of five spots, mainly comprised of mitochondrial proteins including putative mitochondrial HMG-CoA synthase spots, showed a modest induction by lovastatin alone, but little or no effect with any of the other treatments (including the combination of lovestatin and cholestyramine; Fig. 12). This result is intriguing because lovastatin was expected to affect only the regulation of enzymes of cholesterol synthesis, which is entirely extra-mitochondrial. Three of the spots (235, 134, 144) form a closelypacked triad at approximately 30 kDa, and are likely to represent isoforms of one protein. All three spots are stained by an antibody to the mitochondrial form of HMG-CoA synthase obtained from Dr. Greenspan. Subcellular fractionation indicates a mitochondrial location. The other two spots (633 at about 38 kDs and 724 at about 69 kDa) are each present at lower abundance than the members of the triad.

3.3.3 An example of an anti-synergistic effect

A sixth spot (367) shows strong induction by lovastatin (two- to threefold), and about half as much induction with lovastatin plus cholestyramine, but without sharing the animal-animal heterogeneity pattern of the 235-set (Fig. 13). This protein is also mitochondrial, and represents the clearest example of an anti-synergistic effect of lovastatin and cholestyramine. The existence of such an effect demonstrates that lovastatin and cholestyramine do not act exclusively through the same regulatory pathway.

3.3.4 Complexity of the cholesterol synthesis pathway

Taken together, these results suggest that treatment with lovastatin alone can affect both cytosolic and mitochondrial pathways using HMG-CoA, while cholestyramine, on the other hand, either alone or in combination with lovastatin, produces a strong effect on the putative cytosolic pathway, but little or no effect on the putative mitochondrial pathway. An explanation for this difference may lie in lovastatin's effect on levels of HMG-CoA and related precursor compounds that are exchanged between the cytosol and the mitochondrion, whereas cholestyramine should affect only the cytosolic pathways directly controlled by cholesterol and bile acid levels. It remains to be explained why some

proteins of the putative mitochondrial pathway are so much more variable in their expression in all groups. An examination of all the coregulated groups suggests that quantitative statistical techniques can extract a wealth of interesting information from large sets of reproducible gels. The abundance of spots in the 413 coregulation group, for example, shows an amazing level of concordance in their relative expression among the five individuals of the lovastatin and cholestyramine treatment group. This effect is not due to differences in total protein loading, since they have already been removed by scaling, and since proteins with quite different regulation patterns can be demonstrated (e.g., Fig. 13). Such effects raise the possibility that many gene coregulation sets may be revealed through the study of a sufficiently large population of control animals (i.e., without any experimental manipulation). This approach, exploiting natural biological variation in protein expression instead of drug effects, offers an important incentive for the construction of a large library of control animal patterns.

4 Conclusions

Because of the widespread use of rat liver in both basic biochemistry and in toxicology, there is a long-term need for a comprehensive database of liver proteins. The rat liver master pattern presented here has proven to be an accurate representation of this system, having been matched to more than 700 gels to date. As the number of proteins identified and the number of compounds tested for gene expression effects grows, we expect this database to contribute valuable insights into gene regulation. Its practical utility in several areas of mechanistic toxicology is already being demonstrated.

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5 References

- [1] O'Farrell, P., J. Biol. Chem. 1975, 250, 4007-4021.
- [2] Klose, J., Humangenetik 1975, 26, 231-243.
- [3] Scheele, G. A., J. Biol. Chem. 1975, 250, 5375-5385.
- [4] Iborra, G. and Buhler, J. M., Anal. Biochem. 1976, 74, 503-511.
- [5] Anderson, N. G. and Anderson, N. L., Behring. Inst. Mitt. 1979, 63, 169-210.
- [6] Anderson, N. G. and Anderson, N. L., Clin. Chem. 1982, 28, 739-748.
- [7] Heydom, W. E., Creed, G. J. and Jacobowitz, D. M., J. Pharmacol. Exp. Therap. 1984, 229, 622-628.
- [8] Anderson, N. L., Nance, S. L., Tollaksen, S. L., Giere, F. A. and Anderson, N. G., Electrophoresis 1985, 6, 592-599.
- [9] Racine, R. R. and Langley, C. H., Biochem. Genet. 1980, 18, 185-197.
- [10] Klose, J., Mol. Evol. 1982, 18, 315-328.
- [11] Neel, J. V., Baier, L., Hanash, S. and Erickson, R. P., J. Hered. 1985, 76, 314-320.
- [12] Marshall, R. R., Raj, A. S., Grant, F. J. and Heddle, J. A., Can. J. Genet. Cyrol. 1983, 25, 457-446.
- [13] Taylor, J., Anderson, N.L., Anderson, N.G., Gemmell, A., Giometti, C.S., Nance, S. L. and Tollaksen, S. L., in: Dunn, M. J. (Ed.), Electrophoresis '86, Verlag Chemie, Weinheim 1986, pp. 583-587.
- [14] Giometti, C. S., Gemmell, M. A., Nance, S. L., Tollaksen, S. L. and Taylor, J., J. Biol. Chem. 1987, 262, 12764-12767.
- [15] Anderson, N. L., Giere, F. A., Nance, S. L., Gernmell, M. A., Tollaksen, S. L. and Anderson, N. G., in: Galteau, M.-M. and Siest, G. (Eds.), Progrès Récents en Electrophorèse Bidimensionelle, Presses Universitaires de Nancy, Nancy 1986, pp. 253-260.
- [16] Anderson, N. L., Swanson, M., Giere, F. A., Tollaksen, S., Gemmell, A., Nance, S. L. and Anderson, N. G., Electrophoresis 1986, 7, 44-48.

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- [17] Anderson, N. L., Giere, F. A., Nance, S. L., Gemmell, M. A., Tollaksen, S. L. and Anderson, N. G., Fundam, Appl. Toxicol. 1987, 8, 39-50.
- [18] Anderson, N. L., in: New Horizons in Toxicology, Eli Lilly Symposium, 1991, in press.
- [19] Antoine, B., Rahimi-Pour, A., Siest, G., Magdalou, J. and Galteau, M. M., Cell. Biochem. Funct. 1987, 5, 217-231.
- [20] Elliott, B. M., Ramasamy, R., Stonard, M. D. and Spragg, S. P. Biochim. Biophys. Acta 1986, 870, 135-140.
- [21] Huber, B. E., Heilman, C. A., Wirth, P. J., Miller, M. J. and Thorgeirsson, S. S., Hepatology 1986, 6, 209-219.
- [22] Wirth, P. J. and Vesterberg, O., Electrophoresis 1988, 9, 47-53.
- [23] Witzmann, F. A. and Parker, D. N., Toxicol. Lett. 1991, 57, 29-36.
- [24] Rampersaud, A., Waxman, D. J., Ryan, D. E., Levin, W. and Walz, F. G., Jr., Arch. Biochem. Biophys. 1985, 243, 174-183.
- [25] Vlasuk, G. P. and Walz, F. G., Jr., Anal. Biochem. 1980, 105, 112-120.
- [26] Anderson, N. G. and Anderson, N. L., Anal. Biochem. 1978, 85, 331-340.
- [27] Anderson, N. L. and Anderson, N. G., Anal. Biochem. 1978, 85, 341—354.
- [28] Anderson, L., Hofmann, J.-P., Anderson, E., Walker, B. and Anderson, N. G., in: Endler, A. T. and Hanash, S. (Eds.), Two-Dimensional Electrophoresis, VCH Verlagsgesellschaft, Weinheim 1989, pp. 288-297
- [29] Anderson, L., Two-Dimensional Electrophoresis: Operation of the ISO-DALT[®] System. Large Scale Biology Press, Washington, DC 1988, ISBN 0-945532-00-8, 170pp.
- [30] Neuhoff, V., Stamm, R. and Eibl, H., Electrophoresis 1985, 6, 427-448.

- [31] Neuhoff, V., Arold, N., Taube, D. and Ehrhardt, W., Electrophoresis 1988, 9, 255-262.
- [32] Anderson, N. L. and Hickman, B.J., Andl. Biochem. 1979, 93, 312-320.
- [33] Sidman, K. E., George, D. E., Barker, W. C. and Hunt, L. T., Wark. Acids Res. 1988, 16, 1869-1871.
- [34] Taylor, J., Anderson, N.L., Coulter, B.P., Scandora, A. E. and Anderson, N. G., in: Radola, B.J. (Ed.), Electrophoresis 79, de Gruyter, Berlin 1980, pp. 329-339.
- [35] Taylor, J., Anderson, N. L. and Anderson, N. G., in: Allen, R. C. and Arnaud; P. (Eds.), Electrophoresis, 81, de Gruyter, Berlin 1981, pp. 383-400.
- [36] Anderson, N. L., Taylor, J., Scancora, A. E., Coulter, B. P. and Anderson, N. G., Clin. Chem. 1981, 27, 1807-1820, 32
- [37] Taylor, J., Anderson, N. L., Scandora, A. E., Jr., Willard, K. E. and Anderson, N. G., Clin. Chem. 1982, 28, 861–866.
- [38] Taylor, J., Anderson, N. L. and Anderson, N. G., Electrophoresis 1983, 4, 338–345.
- [39] Anderson, N. L. and Taylor, J., in: Proceedings of the Fourth Annual Conference and Exposition of the National Computer Graphics Association, Chicago, June 26–30, 1983, pp. 69–76.
- [40] Anderson, N. L., Hofmann, J.-P., Gemmell, A. and Taylor, J., Clin. Chem. 1984, 30, 2031-2036.
- [41] Anderson, L., in: Schafer-Nielsen, C. (Ed.): Electrophoresis '88, VCH Verlagsgesellschaft, Weinheim 1988, pp. 313-321.
- [42] Neidhardt, F. C., Appleby, D. A., Sankar, P., Hutton, M. E. and Phillips, T. A., Electrophoresis 1989, 10, 116-121.
- [43] Gil, G., Goldstein, J. L., Slaughter, C. A. and Brown, M. S., J. Biol. Chem. 1986, 261, 3710-3716.

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6 Addendum 1: Figures 1–13

Figure 1. Synthetic representation of the standard rat liver 2-D master pattern, rendered as a greyscale image using a videoprinter.

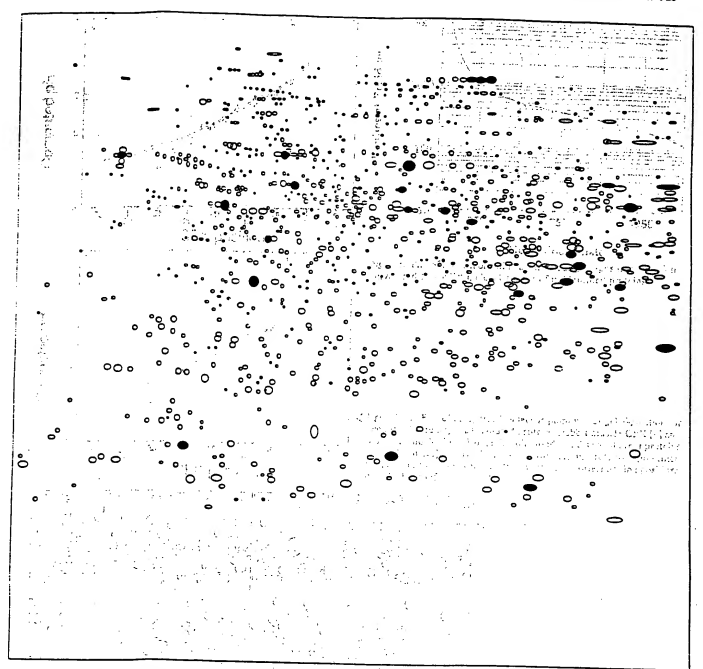


Figure 2. Schematic representation of the master pattern (the same as Fig. 1), useful as an aid in relating specific areas of Fig. 1 and the following detailed quadrants.

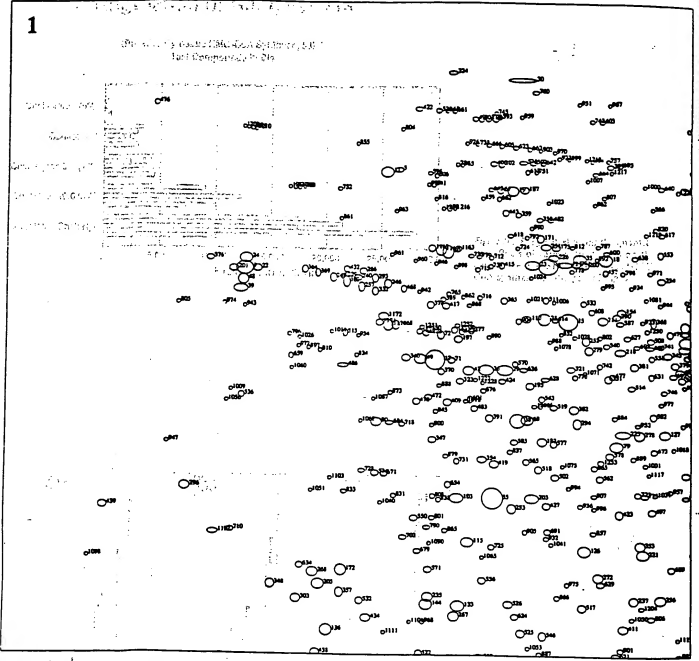


Figure 3. Upper left (high molecular weight, acidic) quadrant (#1) of the rat liver map, showing spot numbers.

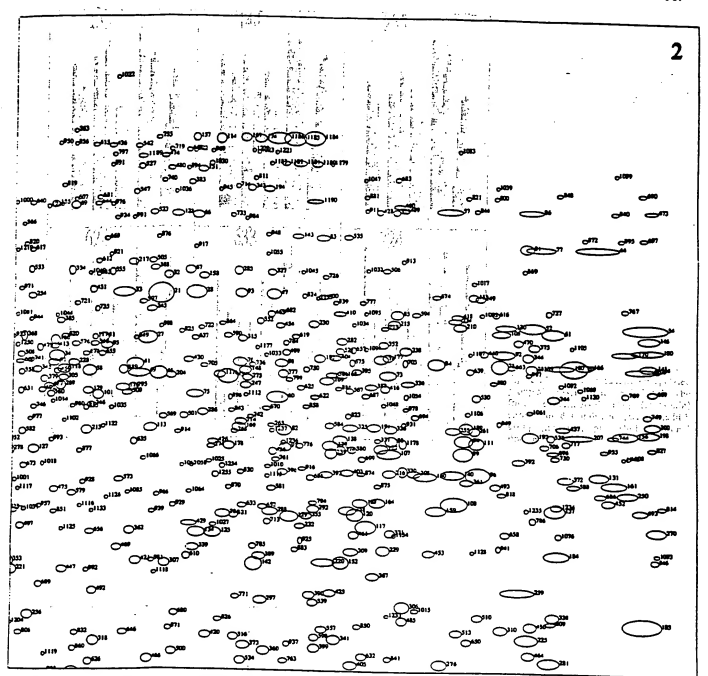


Figure 4. Upper right (high molecular weight, basic) quadrant (#2) of the rat liver map, showing spot numbers.

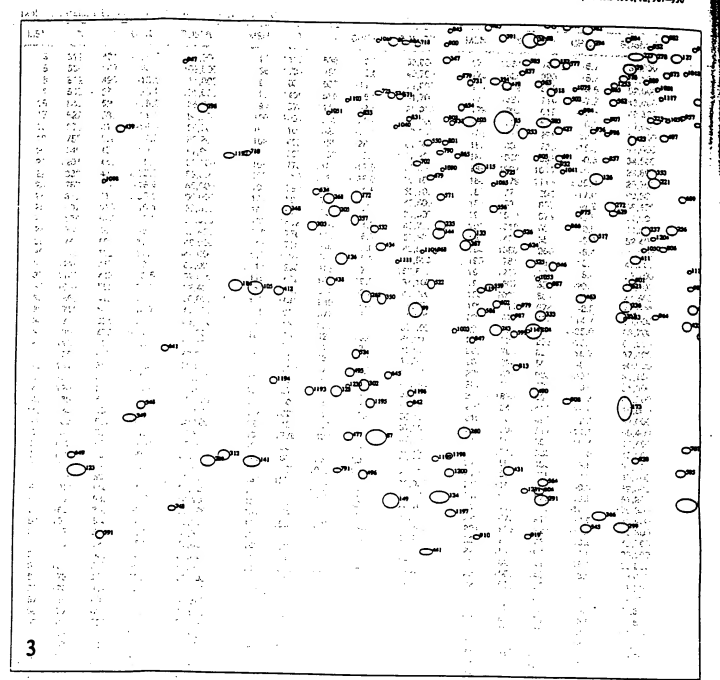


Figure 5. Lower left (low molecular weight, acidic) quadrant (#3) of the rat liver map, showing spot numbers.

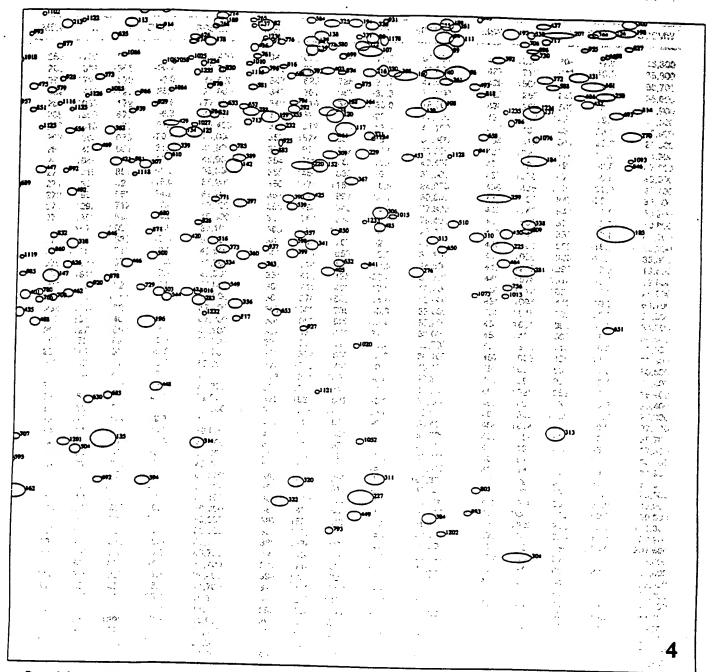


Figure 6. Lower right (low molecular weight, basic) quadrant (#4) of the rat liver map, showing spot numbers.

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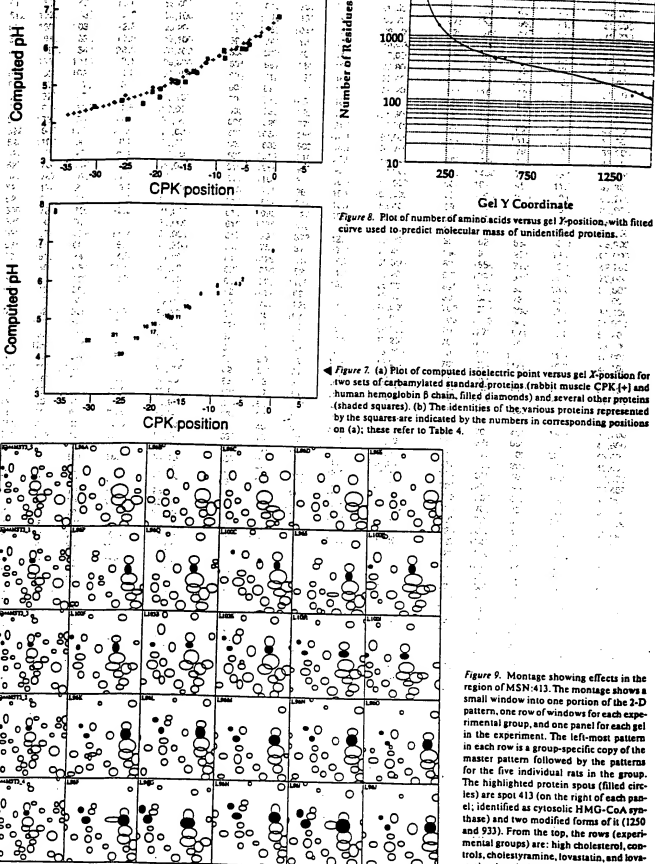
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Figure 9. Montage showing effects in the region of MSN:413. The montage shows a small window into one portion of the 2-D pattern, one row of windows for each experimental group, and one panel for each gel in the experiment. The left-most pattern in each row is a group-specific copy of the master pattern followed by the patterns for the five individual rats in the group. The highlighted protein spots (filled circies) are spot 413 (on the right of each panel; identified as cytosolic HMG-CoA synthase) and two modified forms of it (1250 and 933). From the top, the rows (experimental groups) are: high cholesterol, controls, cholestyramine, lovastatin, and lovastatin plus cholestyramine.

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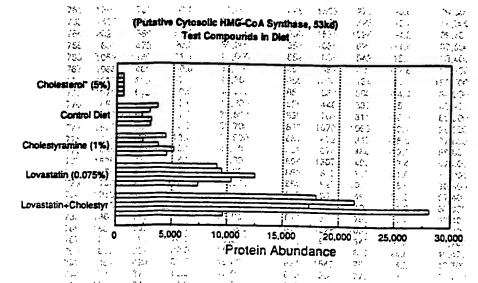
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Regulation of Rat Liver 413



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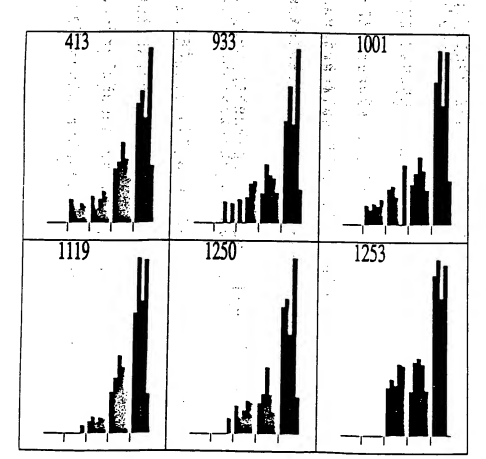


Figure 11. Bargraphs of a series of six coregulated spots including MSN:413. in the bargraphs, the abundances of the appropriate spot (master spot number shown at the top of the panel) in each animal are shown. The five five-animal groups are in . the order (left to right): high cholesterol, controls, cholestyramine, lovastatin, and lovastatin plus cholestyramine. Each bar within a group represents one experimental animal liver (one 2-D gel). Note the correlated expression of the 6 spots, especially in the two far right (most strongly induced) groups.

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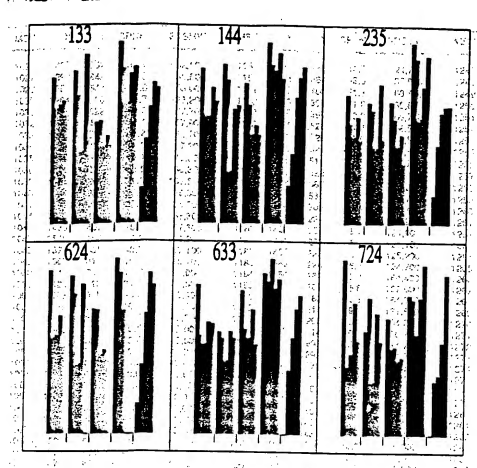


Figure 12. Data on a second coregulated group of spots, presented as in Fig. 11. The fourth experimental group (lovastatin) shows a modest induction, while the fifth group (lovastatin plus cholestyramine) 70 ... 10 ... 11 ... 14 ... 14 ... does not.

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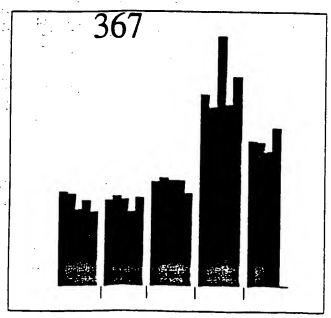


Figure 13. Data on spot MSN:367, presented as in Fig. 11. This protein shows unambiguously the anti-synergistic effect of lovastatin and cholestyramine (fifth group) as compared to lovastatin (fourth group). This response contrasts strongly with the regulation pattern seen in Fig. 11.

Table 1. Master table of proteins in the rat liver database^{a)}

ISN	: X	<i>j</i> : \	CPKol	SDSMW	MSM	ا _, }	(Y	CPKol	SDSMW	MSN	<u>`</u> x	. Y	CPKal	SDSMW
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, ·	1184	524		55,500 54,900	108 109	1692		-2.4	38,300	. 191	1560	674	-3.9	38,400 44,900
ļ	1263	446	-6.0	62,400	110	1482 778	593 516	4.8	49,700	192	1818	687	-0.9	44,200
	743	- 605		49,000	111	1728	700	-16.9 -2.0	55,500 43,500 5	193	1469	555	-5.0	52,400
•	768	₹° 112		348,600	113	1191	680	-8.9	44,500	194	1380 / 784 /	266	-6:4	101,600
	1216 1145	. 417 445		66,000 62,500	114	1298	185	-7.5	160,800	196	1227	632	-16.7 -8:4	23,700
	1037	· 555	-11.3	52,400	115	682	907	-19.6	34,100	197	667	553	-20.1	52,600
	863	412	-14.9	66,600	116 117	1146 1548	610 849	- 9 .5	48,700	198	2006	681	>0.0	44,500
	712	606	-18.7	48,900	148	1050	577	-4.1 -11.1	36,500 50,800	199 200	1711	674.	-2.2	44,900
	763	694	-17.3	43,800	120	1530	828	-4.3	37,400	200	872 292	424	-14.7	65,000
	304 1165	470 569	<-35.0 - 9 .2	59,800 51,400	121	838	423	-15.4	65,200	202	736	253	<-35.0 -18.0	63, 700 107,800
	684	607	-19.6	48,800	122 123	1572 23	712	-3.8	42,900	203	786	829	-16.7	37,400
•	1318	589	-7.3	50,000	124	621	1433 1474	<-35.0 -21.9	15,300	204	1224	589	-8.5	50,000
	1924	362	-0.1	74,600	125	1298	862	-7.5	13,900 36,000	205 206	439 1994	983	-30.9	31,100
	1203	586	-8.7	50,200	126	872	921	-14.7	33,50C	207	1895	571 687	>0.0 -0.3	, 51,300
	1391 309	447 454	-6.3 <-35.0	62,300 61,500	127 128	1000	717	-12.0	42,600	208	240 ^	1418	<-35.0	44,200 15,800
	605	587	-22.5	50,100	129	1229 1422	311 832	-8.4 5:0	86,100	210	1700 _	490	-23	57,000
	621	535	-21.8	53,900	130	1776	499	-5:8 -1.4	37,300 57,000	211 213	902	517	-14.1	55,400
	1113	522	-10.0	55,000	131	1930	757	-0.1	40,700	214	1087 ×	684 668	-10.4	2 44,400
	1820 725 -	499 177	-0.9 -18.3	57,000 ; 170,800	132	660	537	-20:4	53,800	215	1591	495	€7.0 `-3.5	45,200 57,300
:	2001	500	>0.0	56,900	133 134	666 1271	1019	-20.2	29,700	216	1585	755	-3.6	40,700
	722	. 830	-18.4	37,300	135	1161	862 1389	-7.9 - 9 .3	36,000	217	1159	393	-9.3	69,300
	678	533	-19.8	54,100	136	453	1063	-29.7	16,800 28,100	218 219	931 // 713	572 177	-13.5	51,200
	1682 1091	302 580	-2.5 -10.3	89,000 50,600	137	1858	823	-0.6	37,700	220	1479	911	-18.7 -4.9	170,500 33,900
	1171	585	-9.2	50,300	138 139	1504 1488	697	-4.6	43,700	221	965	927	-12.8	33,300
	1400	624	-6.2	47,800	140	1689	707 756	-4.8 -2.4	43,200	223	934	716	-13.5	42,700
	1853	508	-0.6	56,200	141	311	1417	<-35.0	40,700 15,800	225 226	. 1812 5 821	1045	-1.0	28,800
	1888	567	-0.4	51,500	142	1366	915	-6.7	33,800	227	1586	411 1483	-15.8 -3.6	66,800
	735 1 <i>2</i> 63	297 312	-18.1 -8.0	90,500 85,900	143	1429	346	-5.7	77,900	228	1065	567	-10.8	13,600 51,600
	1252	407	-8.1	67,300	144	615 2006	1017	-22.1	29,800	229	1577	890	-3.7	34,800
	779	692	-16.8	43,900	146	2006	566 518	>0.0 >0.0	51,600 55,300	230	1458	496	-5.2	57,300
	1064	296	-10.8	90,800	147	1070	1108	-10.7	26,500	232 234	1440	849 489	-5.5 2.4	36,500
	656 638	589 545	-20.6 -21.2	50,000	148	1347	578	-6.9	50,800	235	618	1004	-2.4 -22.0	57, 900 30, 300
	1582	583	-21.2 -3.6	53,100 50,400	149 150	541 1645	1481	-25.7	13,700	236	920	1138	-13.7	25,400
	1570	556	-3.8		= 151	1269	760 236	-2.8 -7.9	40,500	237	952	1008	-13.1	30,200
	1264	621	-8.0	48,000	152	1507	911	-7.9 - 4 .5	117,000 33,900	238 239	1611 1489	541	-3.2	53,500
	1338 -	564	-7.0	51,800	153	1722	448	-2.1	62,100	240	501	720 448	-4:8 -27.7	42,500
	1833 1767	363 . 565	-0.8 -1.5	74,400 51,700	154	932	503	-13.5	56,600	241	1820	569	-27.7 -0.9	62,100 51,400
	925	738	-1.5 -13.6	51,700 · 41,600	155 156	1031 1970	294 584	-11.4	91,400	242	1357	658	-6.8	45,800
	534	698	-26.1	43,600	157	1258	684 183	>0.0 -8.1	44,400	243	711	1182	-18.7	23,800
	1811	363	-1.0	74,500	158	1275	417	-7.8	162,400 65,900	244 245	1855 1189	621	-0.6	48,000
	1412	681	-6.0 5.0	44,500	159	1663	820	-2.6	37,800	246	551	474 450	-8.9 -25.1	59, 300 61,000
	1471 1662	347 563	-5.0 -2.7	77,500 51,800		1034	527	-11.4	54,600	247	1348	604	-6.9	49,100
	1596	479	-2.7 -3.4	58,900	161 162	1953 1020	771	>0.0	40,000	248	460	448	-29.3	62,100
	817	301	-0.9	89,100		1566	1482 806	-11.6 -3.8	13,700	249	1733	451	-1.9	61,800
	516	1371	-27.0	17,400		1905	565	-3.8 -0.2	38,400 51,700	250 251	1974	788	>0.0	39,200
	589	698	-3.5	43,600	167	1340	181	-7.0	164,900	251 252	808 874	392 553	-16.1 -14.6	69,500
	706	719	-2.2	42,500		1506	583	-4.6	50,400	253	753	333 848	-14.6 -17.6	52, 500 36, 500
	651 415	329 710	-20.8 -6.0	81,700 43,000		1338	678	-7.0	44,700	254	995	450	-12.1	36,500 61, 900
	773	545	-1.4	43,000 53,200	170 171	1969 800	541 370	>0.0	53,500	255	1690	679	.2.4	44,600
	338	446	-7.0	62,300	172	476	378 958	-16.3 -28.7	71,800	256	994	1006	-12.1	30,200
	708	696	-2.2	43.700	173		1314	-28.7 -13.7	32,100	257	508	464	-27.4	60,400

a) Master table of proteins in the rat liver database, showing spot master number, gel position (x and y), isoelectric point relative to CPK standards, and predicted molecular mass (from the standard curve of Fig. 8).

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MSN	la asp	Kalin (S Ý	CPKol	SDSMW	_	MSN	X	Y	СРКЫ	SDSMW	MSN	×	Y	CPKol	SDSMW
259				31,900		345	1006	578	11.9	50,800	426	1296	704	-7.8	49.90-
260		****	-20.4	17,700	?;;		1095		-10.3	46,800	427	810	843	-16.0	43,300 36,800
7, 261 262				25,800 25,800	20	347 348	625		-21.7	42,000	428	1565	303	3.9	88,700
· 263				177,400	5.6		7 361 7 110		-35.3 <-35.0	31,100 1 18,300	429 430	1259	847	-8.0	36,600
€ 265		-	-6.3	45,000	211	350		1130	-26.7	25,700	431	1253 734	562 1426	-8.1 -18.1	51,900
266			-27.3	63,400	4.	351		- ; -	-13.9	48,100	432	483	433	-16.1	15,500 63,900
267 268			-20.4 -31.0	29,000 31,900	ř.		1574	530.	~3.7	54,300	434	518	1041	-26.9	28,900
269			-11.2	48,900	~	353. 354	961 706	912 762	-12.9 -18.9	33,900 40,400	435	1020	1170	-11.6	24,300
270			>0.0	36,300	25	355	1450	830	-5.3	37,300	436 437	1122 1870	196 673	-9.8	147,600
271			-15.0	65,200	3:	356	1374	1152	-6.5	24,900	438	435	1102	-0.5 -31.0	45,000 26,700
., 212			-14.2 -7.6	31,700	구 (원	357		997	-28.7	30,600	439	86	847	<-35.0	36,600
` 274 ' 275			-6.9	42,900 49,900	2. 2.	358°	798 764	346 338	-16.3	77,800 :	440	1740	544	-1.8	53,200
276			-2.6	27,100	24	360	1384	1068	-17.3 -6.4	79,400 <u> </u>	441 443	599	1571	-22.8	10,800
277			-19.4	53,700	22	361	1713	769	-2.1	40,100	446	743 801	335 668	-17.8	80,100
278			-13.0	42,600	11	362	1161	859	-9.3	36,100	447	1050	926	-16.2 -11.1	45,200 33,300
. 279 281	879 1848		-14.5 -0.7	51,300 27,300	*	363	914	1156	-13.8	24,800	448	1245	1298	-8.2	19,800
282			4.6	54,800		364 ·	, 412, 741.	435 486	-32.0	63,700	449	1576	1516	-3.7	12,600
202	1313		-7.3	25,100	2.	366	878	1503	-17.9 -14.6	58,200 13,000	450 451	1818 1094	1021	-0.9	29,600
284	1314	829	-7.3	37,400	95 50	367	1560′	935	-3.9	33,000	452	1945	440 802	-10.3 >0.0	63,100
285 286	1332 1277	408 652	-7.1 -7.8	67,200		368	983	520	-12:4	55,200	453	1652	894	-2.8	38,600 34,600
288	1391	824	-7.5 -6.3	46,100 37,600		369 370	434 639	441 610	-31.0	63,000	454	1403	500	-6.1	56,900
289	1147	579	-9 .5	50,700	ż	371	1587.	860	-21.2 -3.6	48,700 < 36,100 \	456 457	1394	718	-6.3	42,600
<i>2</i> 90	925	511	-13.6	55,900	-	372	1875	762	-0.5	40,400	459	905 1038	436 581	-14.0 -11.3	63,500
291 292	787 1462	1476 818	-16.6 -5.1	13,900		373	1351	1059	-6.8	28,300	460	1598	294	-3.4	50,500 91,400
293	531	449	-26.3	37,800 62,000	8	374 375	1506 1823	715 532	4.6	42,700	461	1528	863	4.3	35,900
294	860	698	-14.9	43,600	•	376	254	417	-0.9 <-35.0	54,200 65,900	462 463	1098	1137	-10.2	25,400
295	1162	609	-0.3	48,700	2	377	1409	583	-6.1	50,400	464	849 1814	1125 1072	-15.2 -0.9	25,800
296 297	218 1377	814 979	<-35.0 -6.5	38,000	ž Ži	378	621	494	-21:8	57,500	465	1388	481	-6.3	27,800 58,700
299	913	1523	-13.9	31 ,300 12, 400		379 381	1017 953	5 95	-11.7	49,600	466	1194	1084	-8.9	27,300
300	2012	667	>0.0	45,300	2	382	856	598 674	-13.1 -15.0	49,400 44,900	468	577	467	-23.9	60,100
301	702	178	-19.0	169,200	٠,	383	1252	258	-8.1	105,300	469 470	1140 1797	888 524	-9.6 -1.1	34,900
302 303	494 403	1280 1006	-28.1 -32.6	20,400 30,100	75.7	384	1699	1518	-2.3	12,500	471	1293	1133	-7.6	54,800 25,500
304	1843	1585	-0.7	10,300	c.r	385 386	1042 1490	493 583	-11.2	57,500	472	618	655	-21.9	46,000
305	1049	503	-11.1	49,800	:	387	1554	603	-4.7 - 4.0	50,400 49,100	473 474	2009 1205	299	>0.0	89,900
306	1608	989	-3.3	30,900	• :	388	1193	404	-8.9	67,700	475	1035	215 788	-8.7 -11.4	131,300 39,200
307 308	1219 1627	916 755	-8.5 3.0	33,700 40,700		389 390	1374	902	-6.5	34,300	476	160	155	<-35.0	207,600
309	1524	892	4.4	34,700		391	1456 718	969 690	-5.2 -18.5	31,700	477	469	1370	-28.9	17,400
310	1769	1028	-1.5	29,400		392	1799	732	-1.1	44,000 41,900	478 479	599 1009	662 540	-22.8	45,600
311	1609	1451	-3.3 - 35.0	14,700		393	1482	758	4.8	40,600	480	1216	235	-11.8 -8.6	53,500 117,400
312 313	266 1902	1408 1365	<-35.0 -0.3	16,100 17,600		394	1227	1461	-8.4	14,400	482	816	346	-15.9	77,800
314	1316	1395	-7.3	16,600		395 - 396	1530 1410	577 755	-4.3 -6.0	50,800 40,800	483	693	673	-19.3	44,900
315	1341	523	-7.0	54,900		397	912	256	-13.9	106,400	485 486	1608 478	1013	-3.3	30,000
318	1104	1053	-10.1	28,500		399	1465	1063	-5.0	28,100	487	1025	5 99 607	-28.6 -11.5	49,300 48,800
320 321	1480 850	1459 603	-4.9 -15.1	14,400 49,100		400 401	1473	450	-4.9	61,900	488	1045	1186	-11.2	23,700
322	1454	1494	-5.3	13,300		401 403	1029 1516	1140 754	-11.5	25,300	· 489	1609	301	-3.3	89,200
323	670	626	-20.0	47,700			1495	554	-4.4 -4.7	40,800 52,500	490 · 491	775 692	1289	-17.0	20,100
324	655	101	-20.6	420,500		405	1525	1092	4.3	27,100	492	1100	178 964	-19.3 -10.2	169,300
325 326	1521 1587	675 677	-4.4 -3.6	44,800 44,700		406	723	252	-18.4	108,000	493	1760	77 6	-10.2	31, 800 39,700
327	1388	409	-6.3	67,000		409 410	650 1501	663 479	-20.8	45,500	494	882	247	-14.5	110,700
328	448	1291	-30.0	20,100		411	936	478 1057	-4.6 -13.4	59,000 28,300	495	470	1258	-28.9	21,200
330	1608	751	-3.3	40,900	4	112	350	1120	-35.9	26,000 26,000	496 497	494 980	1436 852	·28.1	15,200
331 332	1566 531	697 471	-3.8 -26.3	43,700			1033	538	-11.4	53,700	499	1414	546	-12.5 -6.0	36,400 53,100
333	784	1156	-26.3 -16.7	59,600 24,700		115 116	737 1578	425 506	-18.0	64,900	500	1234	1072	-8.3	27,800
334	1059	407	-10.9	67,300		117	646	606 496	-3.7 -21.0	48,900 57,300	501	1246	659	-8.2	45,700
335	1593	303	-3.5	88,500			1695	482	-21.0	57,300 58,600	502 503	824 1246	792 1134	-15.7	39,000
336	1616	598	·3.2	49,400		19	725	770	-18.3	40,000	504	1115	1407	-8.2 -9.9	25,500 16,200
	1854 1265	1004 888	-0.6 -8.0	30,300 34,900				1041	-7.7	28,900	505	1189	391	-8.9	69,700
340	581	585	-23.6	50,300		21 22	1171 5 99	912	-9.1	33,900	506	1578	402	-3 .7	68,000
	1497	1047	₹.7	28,700		23	929	162 856	-22.8 -13.6	193,700 36,200	507 508	787	250	-16.6	109,000
	1351	265		102,200	4	24	739	625	-17.9	36,200 47,700	509	979 1153	552 619	-12.5 -9.4	52,600
344	1813	549	-0.9	52.800	4	25	490	965	4.7	31.800	510	1730	1006	-9.4 -2.0	48,100 30.200
														4.0	J.200

4,350

LICA!		x ·	Y CPKO	SDSMW		J			450.5	_					
MSN		2	· · · · · · ·		MSI	Y X	· '	CPKol			MSN	-	ΥΥ	Jan 1979	SDSWW
511					59						674	91.116 1 661	448	15.27	62,100 %
512 513					56						675	1523		4.4	62,100 ·
514					· · · · · · · · · · · · · · · · · · ·				1.70	25	676		-,	-18.8	46,700
515	48			53,400	600		402				677 678	919 1085		-13.7 -10.5	48,300
516					601		658		• -		679	600		-10.5	52,700 33,400
517 518	86 79		-		602 603		1138				680	1237		-8.3	30,300
519	82			45,100	146.604		181 1461	-14.1 -16.7	165,200	\$ 3	681	1103	283	-10.1	95,100
520	63	•			605		223	-18.0	125,300	4,	682 683	1406 1596	477 249	∕- 6.1	75 7 59,100 7) . ·
521	133	- 1		37,300	606		273	-21.6	98,700		684	555	690	-3.4 -24.8	109,800 43,500
522 523	1190			26,600 86,800	606 608		286			1	685	1167	1313	-9.2	19.300
524	479			22,300	609		503 610	-14.5 >0.0	56,700 48,700	14	686	1932	790	20.0	39,100
525	768	7 + 1		28,000	610		903	-8.1 °	34,200	+ £1	687 688	1545 1456	619 764	-4.1 -5.2	5 48,100
526	747 1170			29,800 L	612		391	-10.1	69,600	1	689	1011	953	-11.8	40,300 4 T
527 528	1502			119,600 53,400	613 614	778 824	265	-16.9	102,000		690	1995	270	>0.0	100,200
530	1728			48,000	615	1095	518 195	-15.7\ -10.3\	55,400 149,100	1	691	812	888	-16.0	34,900
532	507		_	30,000	616	1759	478	-1.6	59,000	51	692 693	1154 1993	1461 819	-9:4 >0.0	14,400 mm
533 534	870 1347			57,900 27,300	617	994	372	-12.1	72,900	£1	694	1628	656	-3.0	45,900
535	1513	346		77,800	618 619	751 1429	374	-17.6	72,400		695	928	254	-13.6	107,000
536	308	654	<-35.0	46,000	620	1050	518 520	-5.7 [:] -11.1.	55,300 1155,200	2) 2)	696	1854	715	-0.6	42,700
538	1851	689		44,100	621	923	1105	-13.7	26,600	21	697 698	1997 ⁻ 957.	345 563	>0.0	78,000
539 540	1463 909	982 561	-5.1 -13.9	31,100 52,000	622 623	1462	622	-5.1	47,900	7.	699	1540	730	-13.0 -4.2	51,800 42,000
541	625	289	-21.7	93,100	624	759 758	225 1038	-17.4 -17.4	124,000		702	577	900	-23:8	34,400 %
542	1164	198	-0 .2	146,200	625	1438	606	-5.5	29,000 48,900	ī -	703. 705	1610. 1278	562	-3.2	51,900
543	803	655	-16.2	45,900	626	1096	1089	-10.2	27,200	,	706	1841	571° 704	-7.8 -0.7	51,200 43,300
544 545	1259 856	1143 1526	-8.0 -15.0	25, 200 12, 200	627 628	942 809	548	-13.3	53,000		707	1018	1386	-11.7	16,900
546	803	1071	-16.2	27,800	629	899	621 979	-16.0 -14.1	48,000 31,300	<u>.</u> 14	709	1074	1145	-10.7	25,100
547	1162	274	.9.3	98,400	630	1135	1321 -	-9.6	19,100		710	_ 293_ _ 720	889 412	<-35.0	34,800
548 549	128 1355	1321 1122	<-35.0 -6.8	19,000 25,900	631	979	615	-12.5	48,300		713	1386	841	-18.5 -6.4	66, 600 36, 800
550	595	866	-23.0	35,800	632 633	1542 1345	1076 814	-4.1 -6.9	27,600		714	1328	263	-7.1	103,100
552	1369	494	-6.6	57,500	634	409	950	-32.2	38,000 32,400		715 716	698 701	433 481	-19.1	63,900
553	992	405	-12.2	67,600	635	1165	704	-9.2	43,300		717	1875	699	-19.0 -0.5	58,700 43,600
555 556	1125 705	410 975	-9.8 -18.9	66,900 31,400	636 637	774 1263	604	-17.0	49,000		718	575	702	-23.9	43,400
557	1477	1030	4.9	29,300	638	952	524 411	-8.0 -13.1	54,800 66,700		719	1216	204	-8.6	140,400
558	980	583	-12.5	50,400	639	1717	575	-2.1	51,000		721 722	1069 1272	464 506	-10.8 -7.9	60,400 56,400
559 560	700 1028	1109 621	-19.1 -11.5	26,400 48,000	640	994	292	-12.1	92,000		723	958	822	-13.0	56,400 37,700
562	898	794	-14.1	38,900	641 642	165 803	1224 251	<-35.0 -16.2	22,400		724	763	395	-17.3	69,100
564	789	1446	-16.6	14,900	643	719	296	-18.5	108,900 90,700		725 726	720 1476	916 415	-18.5	33,700
565 566	777 980	766 328	-16.9	40,200	644	1100	294	-10.2	91,400		727	1846	473	-4.9 -0.7	66,200 59,400
567	1519	611	-12.5 -4.4	81,900 48,600	645 646	534	1263	-26.1	21,000		728	510	783	-27.3	39,400
569	1212	661	-8.6	45,600	648	1153 1246	1038 204	-9.4 -8.2	29,000 140,000		729 730	1217	1126	-8.6	25,800
570	760	594	-17.4	49,700	649	14	1406	<-35.0	16,200		731	1858 665	724 765	-0.6 -20.2	42,300 40,300
571 573	618 1142	956 771	-21.9 -9.6	32,100 40,000	650 651	1713	1049	-2.1	28,600		733	1321	312	-7.2	85, 900
574	532	787	-26.2	39,300	651 652	1986 1378	1183 816	>0.0 -6.6	23,800		734	719	427	-18.5	64,600
575	771	250	-17.1	109,200	653	1442	1165	-6.5 -5.5	38,000 24,400		735 736	1101 1359	473 569	-10.2	59,500
576	1068	534	-10.8	54,100	654	650	806	-20.8	38,400		738	696	220	-6.7 -19.2	51,400 127,600
577 578	822 914	734 754	-15.7 -13.8	41,800 40,800	655 656	1111	551	-10.0	52,700		739	687	409	-19.5	67,000
	1064	794	-10.8	38,900	656 657	1095 1524	861 540	-10.3 -4.4	35,000		740	1205	256	-8 .7	106,200
580	1524	714	4.4	42,800	658	1777	860	-1.4	53,600 36,000		741 742	995 898	563 596	-12.1	51,900
	1392	783	-6.3	39,400	659	391	584	-33.4	50,400		743	881	181	-14.1 -14.5	49,500 165,900
582 584	982 1487	686 672	-12.4 -4.8	44,200 45,000	660 661	977 658	565	-12.5	51,700		744	1951	686	>0.0	44,200
585	758	731	-17.4	41,900	662	732	166 312	-20.5 -18.1	187,500 . 86,100		745 746	726	168	-18.3	183,600
586	687	1152	-19.5	24,900	663	1787	567	-1.2	51,500		748	999 182	643 1503	-12.0 <-35.0	46,600
587 500	930	523	-13.5	55,000	664	888	268	-14,4	100,900		749	2005	649	>0.0	13,000 46,300
588 589	1888 642	774 485	-0.4 -21.1	39,900 58,300	665 666	889	775	-14.3	39,800		750	1448	575	-5.4	51,000
	1317	519	-7.3	55,300	667	715 781	221 227	-18.6 -16.8	126,300		751 752	792	266	-16.5	101,900
591	65	1548	<-35.0	11,500	668	646	165	-10.6	122,400 189,100		75 <u>2</u> 754	469 664	296 254	-28,9 -20.3	90,600
	1014	614	-11.7	48,400	669	1116	353	-9.9	76,300		755	1195	184	-20,3 -8.8	107, 000 161, 000
593 594 1	732 1627	176 478	-18.1 -3.0	172,300 59,000	670 671	1382	643	-6.4	46,600		756	1821	1113	-0.9	26,300
	1009	1426	-11.8	15.500	673	547 984	789 746	-25,3 -12.4	39,200 41.200		757 760	909	246	-13.9	111,000
					-				71.200		, 30	790	133	-16.5	264.900

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MSN	· X	· Y	CPKM	SDSMW	MSN	, X	Y	СРКЫ	SDSMW.	MSN	\ X	Y	CPKal	SDSWW
761 · 763	1399 1416		-5.2 -5.9	41,800 ² 27,300	648 649			-0.6 -6.2	99,500	939		827	-0.0	37,500
754	2020		>0.0	51,400	850			4.2	54,900 29,600	. 1941 942		885 472	-1.5 -22.7	35,000
765	651		-20.8	55,300	851			11.4	37,500	943		496	~35.0	59,600 57,100
766 767	1052 1968		-11.1 >0.0	25,000 59,900	852 855			-15.5	53,400	944		491	-121	57,700
768	: 1330		-7.1	44,300	856		194	-27.8 -10.9	127,100 150,500	945 946	. ,	269 423	-7.5 -21.6	100,300
769	1970		>0.0	48,500	857		890	-14.4	34,800	947	187	736	<35.0	65,100 41,600
770 771	857 1337		-15.0 -7.0	48,200 31,500	858 859	· 1448 706	639 311	-5.4 10.0	46,900	948	1380	344	-6.5	78,200
773	1576		-3.7	56,700	860		1066	-18. 9 -10.7	86,200 28,000	949 950	1766	665	-1.5	
775	969	824	-12.8 -5.5	37,600	861	472	347	-28.8	77,600	951	860	152	-11.3 -14.9	151,000 213,000
776 777	1438 1539	708 . 458	-5.5 -4.2	43,100 61,000	862 864	674 1307	480 499	-19.9 -7.4	58,800	952	957	701	-13.0	43,400
778	850	434	-15.1	63,800	865	645	887	-21.0	57,000 34,900	954 955	503 1938	547, 712	-27.6 >0.0	53,000
779 780	700 1052	411 1136	-19.1 -11.1	66,800 25,500	866	827	1004	-15.6	30,300	957	1010	816	-11.8	42,900 37,900
784	1413	529	-6.0	54,400	968 869	685 1807	494 402	-19.5 -1.0	57,400 68,000	950	768	174	-17.2	174,900
785	1364	885	-6.7	35,000	£70	1323	783	-7.2	39,400	960 961	∴ 596 557	419	-23:0 -24.8	65,700 67,100
786 787	1822 893	835 392	-0.9 ⁽⁺ -14.3	37,100 69,500	671 572	1228	1031	-8.4	29,300	962	887	320	-14.4	83,900
790	616	882	-22.0	35,100	873	1904 556	346 647	-0.3 -24.8	77,700 46,400	963 964	564	334	-24.5	80,500
791	451	1429	-29.8	15,400	874	1540	756	4.2	40,700	965	969 671	1155 255	-12.8 -20.0	24,800 106,600
792 793	777 1536	377 · 1543	-16.9 -4.2	72,000 11,700	875 876	1566 1198	777	-3.8	39,700	966	1204	798	-8.7	38,700
794	1461	807	-5.1	38,300	877	1076	/351 : 720	-8.8 -10.6	76,800 ° 42,500	967 968	910 609	154 1048	-13.9	210,300
196 197	388 1126	546	-33.6 -9.8	53,100	878	1161	1111	-9.3	26,400	969	1285	206	-22.3 -7.7	28,700 138,900
198	933	212 437	-9.6 -13.5	133,700 63,400	879 880	647 1756	757 594	-20.9	40,700	970	822	232	-15.8	119,300
99	1420	593	-5.9	49,800	881	1543	278	-1.6 -4.1	49,700 97,100	971 972	976 403	437 567	-12.6	63,400
00 01	1759 624	279 865	-1.6 -21.7	96,500 35,800	883	1432	890	-5.7	34,800	974	279	495	-32.6 <-35.0	51,600 57;400
02	898	547	-14.2	53,000	884 885	922 1103	689 414	-13.7 -10.1	44,100	975	844	981	15.3	31,200
03	1775	1468	-1.4	14,200	886	1501	607	4.6	66,400 48,900	976 977	1124: 994	295 664	-9.8 -12.1	91,100 45,400
04 05	573 203	196 494	-24.0 <-35.0	148,400 57,400	887 888	798	1103	-16.3	26,600	978	1612	642-	-3.2	46,700
) 6	980	1039	-12.5	29,000	889	636 951	634 759	-21.3 -13.1	.47,200 40,600	979 980	749 1064	1141 642	-17.7	25,300
07 08	902 625	308 827	-14.1 -21.7	87,200	890	717	548	-18.6	52,900	981	1197	911	-10.8 -8.8	46,700 33, 900
9	1851	1015	-0.7	37,500 29,900	891 892	1123 891	229 413	-9.8 -14.3	· 121;200	983	1762	1508	, -1.6	12,800
10	440	573	-30.9	51,100	894	1245	234	-8.2	66,400 117,800	984 985	1344 1024	317 1105	-6.9 -11.5	84,700
1	1358 851	249 393	-6.8 -15.1	109,700 69,400	895 896	1962	346	>0.0	77,700 Ja	987	739	1159	-17.9	26,600 24,600
13	745	1246	-17.8	21,600	897	1322 420	626 570	-7.2 -31.4	47,700 51,300	988 990	816 785	555	-15.9	52,400
14	2028	810	>0.0	38,200	898	662	428	-20.3	64,500	991	1159	361 317	-16.7 - 9 .3	74,900 84,500
15 16	1086 629	645 313	-10.4 -21.6	46,500 85,700	899 900	845 624	243 703	-15.3	113,000	992	1090	928	-10.4	33,300
17	1376	1177	-6.5	24,000	901	931	1094	-21.7 -13.5	43,400 27,000	993	1030 847	701	-11.5	43,400
18 19	1771	790	-1.4	39,100	- 903	799	229	-16.3	121,000	995	902	811 461	-15.2 14.1	38, 200 60,7 00
20	1045 984	263 362	-11.2 -12.4	103,100 74,600	904 905	765 775	520 889	-17.2 -17.0	55,200	996	888	847	-14.4	36,600
21	1712	279	-2.2	96,700	907	888	824	-14.4	34,800 37,600	997 998	1815 1205	579 504	-0.9 -8.7	50,700
23	1256 1517	205 654	-8.1 -4.4	139,200 46,000	906	828	1303	-15.6	19,700	990	617	289	-22.0	56,500 93,100
24	1442	449	-5.5	62,000	910 911	681 1544	1544 301	-19.7 -4.1	11,700 89,100	1000	968	290	-12.8	92,700
25	1240	513	-8.3	55,800	913	1606	387	-3.3	70,400	1001 1002	970 1736	771 478	-12.7 -1.9	40,000 58,900
.6 .7	1309 2012	1014 708	-7.4 >0.0	29,900 43,100	914 916	1237	688	-8.3	44,100	1003	643	1184	-21.1	23,700
18	937	1405	-13.4	16,200	917	1442 1260	749 367	-5.5 -8.0	41,100 73,700	1006 1007	822	487	-15.8	58,100
0	1342	756	-7.0	40,700	919	764	1541	-17.3	11,700	1009	875 291	279 644	-14.6 <-35.0	96,400 46,600
1 2	562 1073	826 1039	-24.5 -10.7	37,500 29,000	920 921	1133	1123	-9.7	25,900	1010	1386	745	-6.4	41,200
3	481	820	-28.5	37,800	923	1123 829	380 242	-9.8 -15.6	71,500 113,200	1011 1012	459	541	-29.4	53,500
4	501	581	-27.8	50,500	924	1131	318	-9.7	84,300	1012	679 1818	661 1128	-19.7 -0.9	45,600 25,800
7 B	751 635	748 833	-17.6 -21.3	41,100 37,200	925 926	1441	874	-5.5	35.400	1014	1032	634	-11.4	47,200
	1494	459	4.7	60,900	926 927	679 1487	219 1191	-19.7 -4.8	128,200 23,500	1015 1016	1629	994	-3.0	30,700
	1952	301	>0.0	89,300	928	1082	775	-10.5	23, 500 39, 800	1016	1311 1722	1134 424	-7.4 -2.0	25,500
1 2	1585 571	1080 1312	-3.6 -24.1	27,500 19,400	929	1231	816	-8.4	38,000	1018	1015	743	-11.7	65,000 41,300
	1325	649	-7.2	46,300	931 932	1609 810	670 900	-3.3 -16.0	45,100 34,400	1020	1574	1219	-3 .7	22,500
4	1727	301	-2.0	89,200	933	965	520	-12.8	34,400 55,100	1021 1022	781 1129	484 83	-16.8 -9.7	58,400 591,300
5 6 :	630 2016	679 905	-21.5 >0.0	44,600 34,200	934	947	462	-13.2	60,600	1023	812	317	-15.9	84,600 ~
		1200	-19.9	23,200	936	865	843	-14.8	36,800	1024	785	446	-16.7	62,400

1028 1030 1031 1032 1033 1034 1035 1036 1039 1041 1045 11051 1052 11053 1056 1058 11066 1067 1068 1069 1061 1065 1066 1067 1068 1069 1061 1073 1075 1076 18 1083 1085 11083 1085 11083 1085 11092 18 1093 1094 15 1095 15 1098 1099 1101 105 18 18 18 18 18 18 18 18 18 18 18 18 18	426	552 848 547 226 822 403 551 496 645 274 262 839 910 485 407 226 635 1040 818 1385 1092 620 377 663 746 605 645 792 839 839 840 840 840 840 840 840 840 840 840 840	-7.7 -12.3 -4.1 -6.4 -4.3 -8.5 -1.6 -25.7 -15.8 -11.3 -5.5 -1.3 -4.2 -31.1 -31.1 -31.1 -31.6 -31.1 -31.6 -31.1 -31.6 -31.1 -31.6 -31	52,600 36,500 53,000 123,200 37,700 67,900 52,700 46,500 98,300 103,600 36,900 34,000 58,300 67,300 109,200 47,100 66,700 28,900 37,800 16,900 48,000 72,000 48,000 41,200 41,200 41,200 41,200 39,000 33,000 33,000 41,800	1153 1154 1162 1163 1168 1168 1170 1171 1172 1174 1176 1177 1178 1179 1180 1181 1183 1184 1185 1186 1189 1190 1191 1192 1193 1194 1195 1195	1594 637 623 665 564 552 538 545 1099	400 397 397 528 529 524 514 522 586 539 702 224 223 223 223 182 214 286 1114 893 1292 1275	-13.7 -3.5 -21.3 -21.8 -20.2 -24.4 -25.9 -25.5 -10.2 -7.5 -6.6 -3.3 -4.8 -5.2 -5.7 -6.1 -6.4 -5.3 -5.8 -6.3 -9.2 -19.5 <-35.0	24,700 35,800 68,400 68,800 68,500 54,500 54,500 55,700 55,000 55,200 55,200 124,900 124,900 125,100 124,900 164,300 164,300 131,800 94,200 26,200 34,700 20,000	1246 1247 1249 1250 1251 1252 1253 1254 1255 1256 1260 1261 1262 1263 1264 1265 1267 1266 1267 1268 1269 1270
1027 1028 1030 1031 1032 1033 1034 1035 1036 1039 1040 1041 1045 1047 1048 1049 1050 1051 1052 1053 1055 1056 1055 1056 1056 1064 11062 1064 1065 1066 1066 1067 1068 1069 1071 1073 1075 1076 1076 1078 1078 1078 1078 1078 1079 1079 1079 1079 1079 1079 1079 1079	1298 1294 986 1547 1381 1525 1128 1226 1761 541 1036 1439 1540 1570 1613 1614 1615 16	848 547 226 822 403 551 496 645 274 262 839 910 485 407 250 635 635 1092 620 377 663 746 604 792 834 758 658 658	-7.5 -15.0 -7.7 -12.3 -4.1 -6.4 -4.3 -9.7 -8.5 -1.6 -25.7 -15.8 -11.3 -5.5 -4.2 -3.7 -10.4 -13.2 -31.1 -3.6 -3.2 -4.5 -3.5 -3.5 -3.6 -3.5 -3.6 -3.5 -3.6 -3.5 -3.6 -3.5 -3.6 -3.5 -3.6 -3.5 -3.6 -3.5 -3.6 -3.6 -3.6 -3.6 -3.6 -3.6 -3.6 -3.6	36,500 53,000 123,200 37,700 67,900 52,700 57,200 46,500 98,300 103,600 36,900 34,000 58,300 67,300 109,200 47,100 27,000 48,000 72,000 48,000 49,000 41,200 49,000 41,200 39,000 33,000	1154 1161 1162 1163 1166 1170 1171 1172 1174 1176 1177 1178 1179 1180 1181 1185 1185 1186 1189 1190 1191 1192 1193 1194 1195 1196	1594 637 623 665 554 552 538 1099 1304 1366 1485 1459 1431 1207 1383 1454 1422 1394 1171 1457 686 265	864 400 397 397 529 524 514 522 586 539 702 224 223 223 224 183 182 214 286 1114 893 1292 1275	-3.5 -21.3 -21.8 -20.2 -24.4 -25.0 -25.9 -25.5 -10.2 -7.5 -6.6 -3.3 -4.8 -5.2 -5.7 -6.1 -5.3 -5.8 -6.3 -9.2 -19.5 -35.0 -32.6	35,900 68,400 68,800 68,700 54,500 54,500 55,700 55,700 55,200 50,200 124,900 124,900 125,100 125,100 126,400 162,600 164,400 162,600 164,300 131,800 94,200 26,200 34,700	1247 1249 1250 1251 1252 1253 1254 1255 1257 1258 1250 1261 1262 1263 1264 1265 1266 1267 1268 1269 1271 1271
1028 1030 1031 1032 1033 1034 1035 1036 1039 1040 1041 1044 1045 1047 1048 1050 1051 1052 11053 1054 11055 1056 1058 11060 1061 1062 11064 11065 1066 1067 1068 1069 1061 1071 1073 1076 1081 1083 1083 1083 1083 1090 1092 18 1093 1093 1093 1093 1093 1093 1093 1093	1284 986 1547 1391 1525 1128 1226 1761 541 618 619 1576 1036 1439 1540 1576 1183 779 1613 1380 284 1261 1281 1285 1275 1181	226 822 403 551 496 645 274 262 839 910 485 407 250 635 411 1040 818 1385 1092 620 377 663 746 605 645 746 792 834 658 696	-7.7 -12.3 -4.1 -6.4 -4.3 -9.7 -8.5 -1.6 -25.7 -15.8 -11.3 -5.5 -4.2 -3.7 -3.6 -16.8 -3.2 -6.5 -3.0 -8.0 -8.0 -8.1 -18.9 -9.0	123,200 37,700 67,900 52,700 57,200 46,500 98,300 103,600 36,900 58,300 67,300 109,200 47,100 66,700 28,900 37,800 16,900 48,000 48,000 41,200 49,000 41,200 39,000 33,000	1162 1163 1166 1170 1171 1172 1174 1176 1177 1178 1189 1180 1181 1185 1186 1189 1190 1191 1192 1193 1194 1195 1196	637 623 665 564 552 538 1099 1304 1366 1608 1485 1459 1431 1407 1383 1454 1422 1394 1171 1457 686 265 403	400 397 397 528 529 524 514 522 586 539 702 224 223 223 223 182 214 286 1114 893 1292 1275	-21.3 -21.8 -20.2 -24.4 -25.0 -25.9 -25.5 -10.2 -7.5 -6.6 -3.3 -4.8 -5.2 -5.7 -6.1 -5.3 -5.8 -6.3 -9.2 -19.5 -35.0 -32.6	68,400 68,800 68,700 54,500 54,500 54,500 55,800 55,200 53,700 43,400 124,900 125,100 125,100 164,400 162,600 164,300 131,800 94,200 26,200 34,700	1249 1250 1251 1252 1253 1255 1257 1258 1259 1260 1261 1262 1263 1264 1265 1266 1267 1268 1269 1270 1271
1031 1032 1033 1034 1035 1036 1039 1040 1041 1044 11044 11047 11048 11049 11050 1051 1052 11053 1054 11055 11056 1058 11060 1061 11062 11061 1066 11066 11067 11068 11060 1061 1067 1068 11060 1061 1067 1068 11060 1061 1067 1068 11060 1061 1071 1075 1076 18 1083 1085 11 1090 1091 1093 1093 1094 11093 1094 11095 11091 1	986 1547 1381 1525 1128 1128 1128 1128 1128 1128 11	822 403 551 496 645 274 262 839 910 485 407 1040 818 1385 1092 620 377 663 746 605 645 746 792 834 658	-12.3 -4.1 -6.4 -4.3 -9.7 -1.6 -25.7 -15.8 -11.3 -5.5 -4.2 -31.1 -3.6 -3.2 -4.5 -3.1 -3.6 -3.2 -4.5 -3.3 -4.9 -8.1 -8.0	37,700 67,900 52,700 57,200 4€,500 98,300 103,600 36,900 34,000 58,300 67,300 109,200 47,100 28,900 37,800 16,900 48,000 48,000 48,000 41,200 49,000 41,200 39,000 33,000	1163 1168 1170 1171 1172 1174 1176 1177 1178 1179 1180 1181 1185 1188 1188 1188 1190 1191 1192 1193 1194 1195 1196	665 564 552 538 545 1099 1304 1366 1608 1485 1451 1407 1383 1454 1472 1394 1477 1457 686 265 403	397 528 529 524 514 522 586 539 702 224 223 224 223 224 182 183 182 214 286 5114 286 1914 1893 1292	-20.2 -24.4 -25.0 -25.9 -25.5 -10.2 -7.5 -6.6 -3.3 -4.8 -5.2 -5.7 -6.1 -6.4 -5.3 -5.8 -6.3 -9.2 -7.5 -6.4 -5.3 -7.5 -7.5 -7.5 -7.5 -7.5 -7.5 -7.5 -7.5	68,800 68,700 54,500 54,500 54,500 55,700 55,700 55,700 124,900 124,900 124,900 125,200 124,700 164,400 162,600 164,300 131,800 94,200 94,200 34,700	1250 1251 1252 1253 1254 1255 1257 1258 1259 1260 1261 1262 1263 1263 1266 1267 1268 1268 1269 1270 1271
1032 1033 1034 1035 1036 1039 1040 1041 1044 11045 11047 11048 11050 1051 1053 1054 11053 1054 11055 11055 11056 11060 1061 11062 11060 1061 11062 11060 1061 11063 11060 11061 11066 11067 11066 11067 11068 11069 11071 1108	1547 1381 1525 1128 1226 1261 541 818 1036 1439 1540 1576 1576 1583 778 1486 1486 1583 778 1486 1486 1583 1617 1245 1258 1705 1181	403 551 496 645 274 262 839 910 485 407 1040 818 1385 1092 620 377 663 746 605 645 746 792 834 658 696	4.1 -6.4 -4.3 -9.7 -8.5 -1.6 -25.7 -15.8 -11.3 -5.5 -2.3 -3.7 -3.6 -16.8 -3.2 -31.1 -3.6 -35.0 -8.0 -33.3 -0.9 -8.1 -18.9 -9.0	67,900 52,700 57,200 46,500 98,300 103,600 36,900 34,000 58,300 67,300 109,200 47,100 65,700 28,900 37,800 16,900 27,000 48,000 48,000 49,000 41,200 39,000 33,000	1168 1170 1171 1172 1174 1176 1177 1178 1179 1180 1181 1182 1183 1184 1185 1190 1191 1192 1193 1194 1195 1195	564 552 538 545 1099 1304 1366 1485 1459 1431 1407 1383 1454 1171 1457 686 265 403 344 505	528 529 524 514 514 522 586 539 702 224 223 224 182 214 286 1114 893 1292 1275	-24.4 -25.0 -25.9 -25.5 -10.2 -7.5 -6.6 -3.3 -4.8 -5.2 -5.7 -6.1 -6.3 -5.8 -6.3 -9.2 -7.5 -8.1 -7.5 -8.1 -7.5 -8.1 -7.5 -8.1 -7.5 -8.1 -7.5 -8.1 -7.5 -8.1 -7.5 -8.1 -7.5 -8.1 -7.5 -8.1 -7.5 -8.1 -7.5 -8.1 -7.5 -8.1 -7.5 -8.1 -7.5 -8.1 -7.5 -8.1 -8.1 -8.1 -8.1 -8.1 -8.1 -8.1 -8.1	54,500 54,500 54,800 55,700 55,700 50,200 53,700 43,400 124,900 125,100 125,100 125,100 164,400 162,600 164,30	1252 1253 1254, 1255 1257 1258 1259 1260 1261 1262 1263 1264 1266 1267 1268 1269 1270 1271 1272
1033 1034 1035 1036 1039 1040 1041 1044 1045 11049 11049 11050 1051 1052 11053 11053 11054 11055 11056 11056 11056 11056 11066 11067 11068 11069 11069 11073 1073	1381 1525 1128 1226 1761 541 541 618 1036 1439 1540 1576 1613 779 426 1583 779 426 1380 284 1261 393 1817 1245 1258 775 1181	551 496 645 274 262 839 910 485 407 255 635 1092 620 377 663 746 605 645 746 792 834 658 696	-6.4 -4.3 -9.7 -8.5 -1.6 -25.7 -15.8 -11.3 -5.5 -4.2 -31.1 -3.6 -16.8 -3.2 -31.1 -3.6 -35.0 -8.0 -33.3 -0.9 -8.1 -18.9 -9.0	52,700 57,200 4E,500 9E,300 103,600 34,000 5E,300 67,300 109,200 47,100 66,700 2E,900 37,800 16,900 27,000 48,000 48,000 49,000 41,200 39,000 39,000 33,000	1170 1171 1172 1174 1176 1177 1178 1179 1180 1181 1183 1184 1185 1186 1189 1190 1191 1192 1193 1194 1195 1196	552 538 545 1099 1304 1366 1608 1485 1459 1431 1407 1383 1454 1171 1457 686 265 403 344 505	529 524 514 522 586 539 702 224 223 223 223 224 182 214 286 1114 893 1292 1275	25.0 -25.9 -25.5 -10.2 -7.5 -6.6 -3.3 -4.8 -5.2 -5.7 -6.1 -6.4 -5.3 -5.8 -6.3 -9.2 -5.2 -19.5 -35.0 -32.6	54,500 54,800 55,700 55,000 50,200 53,700 43,400 124,900 125,100 125,100 126,200 164,400 162,600 164,300	1253 1254, 1255 1257 1258 1259 1260 1261 1262 1263 1264 1265 1266 1267 1268 1269 1270 1271 1272
1035 1036 1039 1040 1041 1044 1 1045 1 1047 1 1048 1 1053 1054 1 1055 1 1056	1128 1226 1761 541 818 1036 1439 1540 1576 1613 1613 1613 1920 1921 1921 1921 1921 1921 1921 1921	645 274 262 839 910 485 407 250 635 411 1048 1385 1092 620 377 663 746 605 645 746 792 934 658 696	-9.7 -8.5 -1.6 -25.7 -11.3 -5.5 -4.2 -31.1 -3.6 -13.2 -31.1 -3.6 -3.2 -6.5 -3.2 -6.0 -33.3 -0.9 -8.1 -8.0 -9.0	46,500 98,300 103,600 36,900 34,000 58,300 67,300 106,200 47,100 65,700 28,900 37,800 16,900 27,000 48,000 49,000 41,200 49,000 38,000 33,000	1172 1174 1176 1177 1178 1179 1180 1181 1185 1185 1186 1189 1190 1191 1192 1193 1194 1195 1196	545 1099 1304 1366 1485 1485 1431 1207 1383 1454 1422 1394 1171 1457 686 265 263 344 505	524 514 522 586 539 702 224 223 223 223 224 182 214 286 1114 893 1292 1275	25.9 -25.5 -10.2 -7.5 -6.6 -3.3 -4.8 -5.2 -5.7 -6.1 -6.4 -5.3 -6.3 -9.2 -19.5 <35.0	54,800 55,700 55,000 50,200 53,700 43,400 124,900 125,100 125,100 126,400 164,400 162,600 164,300 131,800 94,200 26,200 34,700	1254 1255 1257 1258 1259 1260 1261 1262 1263 1263 1265 1266 1267 1268 1269 1270 1271 1272
1036 1039 1040 1041 1046 1047 1048 1049 1050 1051 1053 1054 1053 1054 1055 1056 1056 1066 1067 1068 1066 1067 1068 1066 1067 1068 1069 1071 1073 1075 1081 1083 1085 1081 1083 1083 1085 1081 1083 1083 1083 1083 1083 1083 1083 1083 1083 1083 1083 1083 1094 1095 1092 18 1093 1094 1095 1095 1096 1091 1093 1094 1095 1095 1096 1097 1098 1099 1091 1093 1094 1095 1096 1097 1098 1099 1091 1093 1094 1095 1096 1097 1098 1099 1091 1093 1094 1095 1096 1097 1098 1099 1091 1093 1094 1095 1096 1097 1098 1099 1091 1093 1094 1095 1096 1097 1098 1099 1091 1093 1094 1095 1096 1097 1098 1099 1091 1093 1094 1095 1096 1097 1098 1099 1091 1093 1094 1095 1096 1097 1098 1099 1091 1093 1094 1095 1096 1097 1098 1099 1091 1091 1092 1093 1094 1095 1096 1097 1098 1099 1091 10	1226 1761 541 818 1036 1439 1540 1576 1089 649 426 1583 779 1613 1380 284 1261 284 1261 1245 1245 1705 1181	274 262 839 910 485 407 250 635 411 1040 818 1385 1092 620 377 663 746 605 645 746 792 934 734 658 696	-8.5 -1.6 -25.7 -15.8 -11.3 -5.5 -4.2 -3.7 -10.4 -13.2 -31.1 -3.6 -16.8 -3.2 -4.5 <35.0 -8.0 -33.3 -0.9 -8.1 -18.9 -9.0	98,300 103,600 36,900 34,000 58,300 67,300 109,200 47,100 66,700 28,900 37,800 16,900 27,000 48,000 41,200 49,000 41,200 39,000 33,000	1174 1176 1177 1178 1179 1180 1181 1182 1183 1184 1185 1186 1189 1190 1191 1192 1193 1194 1195 1196	1099 1304 1366 1608 1485 1431 1407 1383 1454 1422 1394 1171 1457 686 265 403	522 586 539 702 224 223 223 224 182 182 214 286 1114 893 1292 1275	-10.2 -7.5 -6.6 -3.3 -4.8 -5.2 -5.7 -6.1 -6.4 -5.3 -6.3 -9.2 -5.9 -35.0 -32.6	55,700 55,000 50,200 53,700 43,400 124,900 125,100 125,200 124,700 164,400 162,600 161,800 94,200 94,200 34,700	1255 1257 1258 1259 1260 1261 1262 1263 1264 1265 1266 1267 1268 1269 1270 1271 1272
1039 1040 1041 1044 11 1044 11 1045 11 1049 11 1050 1051 1055 11 1055 11 1055 11 1056 11 1066 11 1067 1068 11 1071 1073 1075 18 1083 1085 11 1090 1091 1092 18 1099 19 11 1093 1094 11 1095 15 1098 11 1095 15 1098 11 1095 15 1098 11 1091 1091 1091 1091 11 11 11 11 11 11 11 11 11 11 11 11 1	1761 541 818 1036 1439 1540 1576 1089 949 426 1583 779 1181 1380 294 1261 393 1817 1245 1245 1705 1181	262 839 910 485 407 250 635 411 1040 818 1385 1092 620 377 663 746 605 645 746 792 834 734 658 696	-1.6 -25.7 -15.8 -11.3 -5.5 -4.2 -3.7 -10.4 -13.2 -31.1 -3.6 -3.2 -6.5 <35.0 -8.0 -33.3 -0.9 -8.1 -18.9 -9.0	103,600 36,900 34,000 58,300 67,300 105,200 47,100 66,700 28,900 16,900 27,000 48,000 72,000 48,000 41,200 46,600 41,200 33,000 33,000	1176 1177 1178 1179 1180 1181 1182 1183 1184 1185 1186 1189 1190 1191 1192 1193 1194 1195 1196	1304 1366 1608 1485 1459 1431 1407 1383 1454 1422 1394 1171 1457 686 265 403 404 505	586 539 702 224 223 223 223 224 182 214 286 1114 893 1292 1275	-7.5 -6.6 -3.3 -4.8 -5.2 -5.7 -6.4 -5.3 -5.8 -6.3 -9.2 -19.5 -35.0	50,200 53,700 43,400 124,900 125,100 125,100 125,200 164,400 162,600 164,300 131,800 94,200 26,200 34,700	1258 1259 1260 1261 1262 1263 1264 1265 1266 1267 1268 1269 1270 1271 1272
1040 1041 1044 1045 1047 1048 1049 1050 1051 1053 1054 1055 1056 1058 1056 1066 1061 1062 1066 1067 1068 1066 1067 1068 1069 1075 1076 1075 1076 1075 1076 1071 1075 1076 1077 1078 1079	818 1036 1439 1540 1576 1089 949 426 1583 779 1613 1380 284 1261 393 1817 1245 1258 705 1181	910 485 407 255 411 11040 818 1385 1095 620 377 663 746 605 645 746 792 934 734 658 696	-15.8 -11.3 -5.5 -3.7 -10.4 -13.2 -31.1 -3.6 -16.8 -3.2 -6.5 -35.0 -8.0 -33.3 -0.9 -8.2 -8.1 -18.9 -9.0	34,000 58,300 67,300 109,200 47,100 65,700 28,900 37,800 16,900 27,000 48,000 45,500 41,200 49,000 46,600 41,200 39,000 33,000	1177 1178 1179 1180 1181 1182 1183 1184 1185 1190 1191 1192 1193 1194 1195 1196	1366 1608 1485 1485 1497 1383 1454 1422 1394 1171 1457 686 265 403 344 505	539 702 224 224 223 223 224 182 183 182 214 286 1114 893 1292 1275	-6.6 -3.3 -4.8 -5.2 -5.7 -6.4 -5.3 -5.8 -6.3 -9.2 -5.2 -19.5 -35.0	53,700 43,400 124,900 125,100 125,100 125,200 164,400 162,600 164,300 131,800 94,200 26,200 34,700	1259 1260 1261 1263 1263 1264 1265 1266 1267 1268 1269 1270 1271 1272
1044 1 1045 1 1047 1 1048 1 1049 1 1050 1 1051 1 1052 1 1053 1 1056 1 1056 1 1056 1 1066 1 1067 1 1068 1 1067 1 1068 1 1067 1 1073 1 1073 1 1076 1 1078 8 1081 6 1078 8 1081 1 1079 1 10	1036 1439 1540 1576 1089 949 426 1583 779 1613 1380 284 1261 393 1817 1245 1258 705 1181	485 407 250 635 11040 818 1385 1092 620 377 663 746 605 645 746 792 934 658 696	-11.3 -5.5 -4.2 -3.7 -10.4 -13.2 -31.1 -3.6 -16.8 -3.2 -6.5 -35.0 -8.0 -33.3 -0.9 -8.1 -18.9 -9.0	58,300 67,300 109,200 47,100 66,700 28,900 37,800 16,900 27,000 48,000 45,500 41,200 49,000 41,200 39,000 33,000	1179 1180 1181 1182 1183 1184 1189 1190 1191 1192 1193 1194 1195 1196	1485 1459 1431 1207 1383 1454 1422 1394 1171 1457 686 265 403 344 505	224 224 223 223 224 182 183 182 214 286 1114 893 1292 1275	-3.3 -4.8 -5.2 -5.7 -6.4 -5.3 -5.8 -6.3 -9.2 -5.2 -19.5 <-35.0	43,400 124,900 124,900 125,100 125,200 124,700 164,400 162,600 164,300 164,300 164,300 94,200 94,200 34,700	1260 1261 1262 1263 1265 1266 1266 1267 1268 1269 1270 1271 1272
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1047 1 1048 1 1049 1 1050 1 1051 1 1052 1 1053 1 1054 1 1055 1 1056 1 1066 1 1067 1 1068 1 1066 1 1067 1 1071 1 1073 1 1075 1 1076 1 1077 1 1078 1 1083 1 1083 1 1083 1 1083 1 1083 1 1094 1 1095 1 1098 1 1099 1 1093 2 1094 1 1095 1 1098 1 1099 1 1091 1 1093 1 1095 1 1098 1 1099 1 1091 1 10	1540 1576 1089 949 426 1583 779 1613 1380 284 1261 393 1817 1245 1258 705 1181	250 635 411 1040 1385 1092 620 377 663 746 605 645 746 792 934 658 696	4.2 3.7 10.4 13.2 -31.1 3.6 16.8 3.2 -6.5 < 35.0 -8.0 33.3 -0.9 -8.1 -18.9 -9.0	109,200 47,100 66,700 28,900 37,800 16,800 27,000 48,000 72,000 45,500 41,200 46,600 41,200 39,000 33,000	1181 1182 1183 1184 1185 1186 1189 1190 1191 1192 1193 1194 1195 1196	1431 1407 1383 1454 1422 1394 1171 1457 686 265 403 344 505	223 223 224 182 183 182 214 286 1114 893 1292 1275	-5.7 -6.4 -5.3 -5.8 -6.3 -9.2 -5.2 -19.5 <-35.0	125,100 125,200 124,700 164,400 164,500 164,300 131,800 94,200 26,200 34,700	1263 1264 1265 1266 1267 1268 1269 1270 1271 1272
1049 1 1050 1 1051 1 1052 1 1053 1 1054 1 1055 1 1056 1 1056 1 1066 1 1061 1 1062 1 1066 1 1067 1 1068 1 1069 1 1071 6 1071 6 1071 6 1078 8 1081 1 1075 1 1076 1 1078 1 1079 1 10	1089 549 426 1583 779 1613 1380 284 1261 393 1817 1245 1258 705 1181	411 1040 818 1385 1092 620 377 663 746 605 645 746 792 934 658 658	-10.4 -13.2 -31.1 -3.6 -16.8 -3.2 -6.5 -35.0 -8.0 -33.3 -0.9 -8.2 -8.1 -18.9 -9.0	65,700 28,900 37,800 16,900 27,000 48,000 72,000 45,500 41,200 46,600 41,200 39,000 33,000	1183 1184 1185 1186 1189 1190 1191 1192 1193 1194 1195 1196	1407 1383 1454 1422 1394 1171 1457 686 265 403 344 505	223 224 182 183 182 214 286 1114 893 1292 1275	-6.1 -6.4 -5.3 -5.8 -6.3 -9.2 -5.2 -19.5 <-35.0	125,200 124,700 164,400 162,600 161,800 131,800 94,200 26,200 34,700	1264 1265 1266 1267 1268 1269 1270 1271 1272
1050 1051 1052 1053 1054 1056 1058 1056 1056 1066 1061 1062 1064 1065 1066 1067 1068 1069 1071 1073 1075 1076 1078 1079 1083 1083 1093 1093 1093 1094 1093 1094 1095 1095 1096 1093 1099 1093 1094 1095 1096 1097 1098 1099	\$49 426 1583 779 1613 1380 284 1261 393 1817 1245 1258 705 1181	1040 818 1385 1092 620 377 663 746 605 645 746 792 934 658 696	-13.2 -31.1 -3.6 -16.8 -3.2 -6.5 -35.0 -8.0 -33.3 -0.9 -8.2 -8.1 -18.9 -9.0	28,900 37,800 16,900 27,000 48,000 45,500 41,200 49,000 46,600 41,200 39,000 33,000	1184 1185 1186 1189 1190 1191 1192 1193 1194 1195	1454 1422 1394 1171 1457 686 265 403 4344 505	182 183 182 214 286 1114 893 1292 1275	-5.3 -5.8 -6.3 -9.2 -5.2 -19.5 <-35.0	124,700 164,400 162,600 164,300 131,800 94,200 26,200 34,700	1265 1266 1267 1268 1269 1270 1271 1272
1051	426 1583 779 1613 1380 284 1261 393 1817 1245 1258 705 1181	818 1385 1092 620 377 663 746 605 645 746 792 834 734 658 696	-31.1 -3.6 -16.8 -3.2 -6.5 -35.0 -8.0 -33.3 -0.9 -8.2 -8.1 -18.9 -9.0	37,800 16,800 27,000 48,000 72,000 45,500 41,200 49,000 46,600 41,200 39,000 33,000	1185 1186 1189 1190 1191 1192 1193 1194 1195 1196	1422 1394 1171 1457 686 265 403 4344 505	183 182 214 286 1114 893 1292 1275	-5.8 -6.3 -9.2 -5.2 -19.5 <-35.0	162,600 164,300 131,800 94,200 26,200 34,700	1267 1268 1269 1270 1271 1272
1053 1054 1055 1056 1058 1058 1060 1061 1062 1064 1065 1066 1067 1068 1069 1071 1073 1075 1075 1076 1077 1078 1079 1083 1083 1083 1083 1083 1083 1083 1083 1090 1092 18 1093 1094 11095 15 1098 11095 15 1098 11095 15 1098 1099 1091 1092 1093 1094 1095 1096 1097 1098 1099 1091 1093 1094 1095 1096 1091 1091 1093 1094 1095 1096 1097 1098 1099 1091 1091 1093 1094 1095 1096 1097 1098 1099 1099 1091	779 1613 1380 284 1261 393 1817 1245 1258 705 1181	1092 620 377 663 746 605 645 746 792 934 734 658 696	-16.8 -3.2 -6.5 <-35.0 -8.0 -33.3 -0.9 -8.2 -8.1 -18.9 -9.0	.27,000 48,000 72,000 45,500 41,200 49,000 46,600 41,200 39,000 33,000	1186 1189 1190 1191 1192 1193 1194 1195 1196	1394 1171 1457 686 265 403 344 505	182 214 286 1114 893 1292 1275	-6.3 -9.2 -5.2 -19.5 <-35.0 -32.6	164,300 131,800 94,200 26,200 34,700	1268 1269 1270 1271 1272
1054 1 1055 1 1056 1 1058 1 1060 1 1061 1 1062 1 1064 1 1065 1 1066 1 1067 1 1067 1 1075 1 1076 1 1077 1 1078 1 1079 1 1079 1 1081 1 1092 1 1093 2 1094 1 1095 1 1099 1 1099 1 1099 1 1099 1 1099 1 1090 1 1091 1 1095 1 1095 1 1096 1 1097 1 1098 1 1099 1 1091 1 1093 1 1094 1 1095 1 1096 1 1097 1 1098 1 1099 1 1091 1	1613 1380 284 1261 393 1817 1245 1258 705	620 377 663 746 605 645 746 792 934 734 658 696	-3.2 -6.5 -35.0 -8.0 -33.3 -0.9 -8.2 -8.1 -18.9 -9.0	48,000 72,000 45,500 41,200 49,000 46,600 41,200 39,000 33,000	1190 1191 1192 1193 1194 1195 1196	.1457 686 265 403 :344 505	286 1114 893 1292 1275	-9.2 -5.2 -19.5 <-35.0	131,800 94,200 26,200 34,700	1269 1270 1271 1272
1055 1 1056 1 1058 1 1060 1 1061 1 1062 1 1064 1 1065 1 1066 1 1067 1 1068 1 1071 1 1071 1 1075 1 1076 1 1078 1 1079 1 1093 1 1093 1 1093 1 1094 1 1095 1 1098 1 1099 1 1091 1 1093 1 1094 1 1095 1 1096 1 1097 1 1098 1 1099 1 1091 1 10	1380 284 1261 393 1817 1245 1258 705	377 663 746 605 645 746 792 934 734 658 696	-6.5 <-35.0 -8.0 -33.3 -0.9 -8.2 -8.1 -18.9 -9.0	72,000 45,500 41,200 49,000 46,600 41,200 39,000 33,000	1191 1192 1193 1194 1195 1196	686 265 403 ::344 505	1114 893 1292 1275	-19.5 <-35.0 32.6	26,200 34,700	1271 1272
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1060 1061 1 1062 1 1065 1066 1 1067 1 1067 1 1073 1 1075 1 1075 1 1076 1 1078 1 1078 1 1083 1 1083 1 1085 1 1092 1 1093 1 1094 1 1095 1 1 1 1 1 1 1 1 1	393 1817 1245 1258 705 1181	605 645 746 792 934 .734 658 696	-33.3 -0.9 -8.2 -8.1 -18.9 -9.0	49,000 46,600 41,200 39,000 33,000	1194 1195 1196	±.344 = 505	1292 1275	32.6	-	
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1065	705 1181	934 734 658 696	-18.9 -9.0	33,000	1197		1311 1293	-27.6 -24.1	19,400 20,000	1277
1066	1181	734 658 696	-9.0			639	1502	-21.2	13,000	1278 1279
1067 1 1068 1 1069 1 1071 1 1073 1 1075 8 1076 18 1078 8 1083 16 1085 1 1090 6 1092 18 1093 20 1094 15 1098 1 1099 1 1101 5 1102 10 1103 1 1105 18 11105 18 11106 17 11107 17 1108 19 1111 5 11115 13 1115 13 1117 9 1118 12 1119 103		658 696			1198	637	1402	-21.3	16,300	1280
1069 11 1071 6 1071 6 1073 17 1075 8 1078 8 1081 8 1083 16 1085 11 1090 18 1093 20 1094 15 1098 15 1098 15 1099 19 1101 5 1102 10 1103 4 1105 18 11106 17 1106 17 1108 19 1111 5 1111 5 11111 5 11111 15 1111 15 111 15 11 15 11 15 11 15 1				.45,800	1199 1200	614 637	1407 1431	-22.1 -21.3	16,200	1281
1071	508		-27.4	43,700	1201	1095	1394	-10.3	15,400 16,600	1282 1283
1073 17 1075 8 1076 18 1076 18 1081 9 1083 16 1085 11 1090 6 1093 20 1094 15 1095 15 1098 1099 19 1101 5 1102 10 1103 4 1105 18 1106 17 1107 17 1108 19 1111 5 1112 13 1115 13 1115 13 1115 13 1115 13 1117 9 1118 12	873	604 (.	-0.3 -14.7	49,100 48,700	1202	1719	1545	-2.1	11,600	1284
1076 18 1078 8 1081 6 1083 16 1083 16 1090 6 1092 18 1093 20 1094 15 1098 1 1099 19 1101 5 11102 10 1102 10 1103 4 11105 18 11106 17 11108 19 1111 5 11112 13 1115 13 1115 13 1115 13 1117 9 1118 120	768	1128	-1.5	25,800	1203 1204	791 964	668 1021	-16.5 -12.9	45,200	1285
1078	836	773	-15.4	39,900	1205	313	195	<-35.0	29,700 148,700	. 1286 1287
1081	863 / 826 /	961 566	-0.6 -′-15.7	36,000 51,600	1208 1209	306	194	<-35.0	149,800	1288
1085 11 1090 6 1092 18 1093 20 1094 15 1095 15 1098 1099 19 1101 5 1102 10 1103 4 1105 18 1106 17 1107 17 1108 19 1111 5 11112 13 1115 13 1115 13 1115 13 1117 9 1118 12	971	483	-12.7	58,500	1210	320 326	197 197	<-35.0 <-35.0	147,400 146,600	1289 . 1290
1090 6 1092 18 1093 20 1094 15 1095 15 1098 1099 19 1101 5 1102 10 1103 4 11105 18 1106 17 1108 19 1111 5 11112 13 11115 13 11116 10 1117 9 1118 12 1119 10	697	202	-2.3	142,300	1211	394	294	-33.2	91,400	1291
1092 18 1093 20 1094 15 1095 15 1098 19 1101 5 1102 10 1103 4 1105 18 1106 17 1108 19 1111 5 11112 13 1115 13 1115 13 1116 10 1117 9 1118 12 1119 10	157 620	.910	-9.4 -21.9	38,900 34,000	1212	402	294	-32.7	91,200	1292
1094 15 1095 15 1098 19 1109 19 1101 5 1102 10 1103 4 1105 18 1106 17 1107 17 1108 19 1111 5 11112 13 1115 13 1115 13 1115 13 11116 10 1117 9 1118 12 1119 10	867	5 0 7	-0.5	49,500	1214 1215	386 641	294 329	-33.7 -21.2	91,400	1293
1095 15 1098 1 1099 19 1100 5 1100 10 1102 10 1105 18 1106 17 1107 17 1108 19 1111 5 11112 13 1115 13 1115 13 1117 9 1118 12 1119 10	019	894	>0.0	34,600	1216	660	329	-20.4	81,600 81,600	1294 1295
1098 1099 19 1101 5 1102 10 1103 4 1105 18 1106 17 1107 17 1108 19 1111 5 1112 13 1115 13 1116 10 1117 9 1118 12 1119 10		538 477	. च.१ च.१ %	53,700	1217	914	266	-13.8	101,800	
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1102 10 1103 4 1105 18 1106 17 1106 17 1108 19 1111 5 1112 13 1115 13 1115 10 1117 9 1118 12 1119 10	954	237	>0.0	116,000	1220	1021	298	-12.7 -11.6	72,900 90,100	
1103 4 1105 18 1106 17 1107 17 1108 19 1111 5 11112 13 11115 13 11116 10 1117 9 1118 12 1119 10	588 -	1048	-23.3	28,600	1221	1392	205	-6.3	139,500	
1105 18 1106 17 1107 17 1108 19 1111 5 11112 13 11115 13 11116 10 1117 9 1118 12 1119 10	457	667 797	-11.1 -29.5	45,200 38,800	1222 1223	1354 1362	203 205	-6.8	141,800	
1107 17 1108 19 1111 5 1112 13 1115 13 1116 10 1117 9 1118 120 1119 10	884	532	-0.4	54,200	1224	673	540	-6.7 -19.9	139,500 53,600	
1108 19 1111 5 1112 13 1115 13 1116 10 1117 9 1118 120 1119 10	714	649	-2.1	46,300	1225	614	542	-22.1	53,400	
1111 5 1112 13 1115 13 1116 10 1117 9 1118 12 1119 10		546 722	-2.1 >0.0	53,100 42,400	1226 1227	603 696	539	-22.6	53,600	
1115 136 1116 10 1117 9 1118 126 1118 10	547	1066	-25.3	28,000	1228	707	623 628	-19,2 -18,9	47,800 47,500	
1116 10 1117 9 1118 120 1119 10	348	621	-6.9	48,000	1229	475	447	-28.7	62,300	
1117 91 1118 120 1119 100		762 816	-6.4 -10.6	40,400 38,000	1230	466	1282	-29.0	20,400	
1118 120 1119 10		787	-12.6	39,300	1231 1232	759 1324	1461 1170	-17.4	14,400	
	75	933	-8.7	33,100	1233	1583	1005	-7.2 -3.6	24,200 30,300	
ייי חבוו	202	1076	-11.6	27,600	1234	1865	809	-0.6	38,200	
	202 202	616 1301	-0.3 -4.5	48,300 19,700	1235 1236	1812 1411	817	-1.0	37,900	
1122 111	202 222 205	677	-9.9	44,700	1236	1392	703 682	-6.0 -6.3	43,400 44,500	
1123 146	202 022 005 512	452	-5.1	61,700	1238	794	410	-16.4	66,900	
	202 022 005 512 114	857	-11.1 -9.8	36,200 38,600	1239	769	407	-17.1	67,300	
1128 172	202 022 005 512 114 164 148		- 2 .1	34,700	1240 1241	740 743	406 511	-17.9 -17.9	67,500 55,900	
133 109	202 022 005 512 114 164 148 22	802 892		37,500	1242	713	510	-17.8 -18.7	55,900 56,000	•
139 183	202 022 005 512 114 164 148 22 22	802 892 825	-10.2	51,400	1243	682	509	-19.6	56,100	
147 76 148 196	202 022 005 512 114 164 148 22 122 98	802 892	-10.2 -0.8 -17.3	23,800	1244	663	504	-20.3	56,500	

		Databa	res of tat	liver protei	1
•	MSN	×	- Y	СРКЫ	SDSANN
	-	1, 1	C +1750	1 1	
	1246	547	577	-25.3	50 000
	1247	530	576_	-26.3	50,800
	1249	516	572	-27.0	50,900
	1250	973	.536	12.7	51,200
•	1251	607	532	-22.4	53,900 (%)
	1252	665	529	-20.2	54,200 G
	1253	899	766	-14.1	40,200
	1254,	1311	746,		41,200
	1255	1300	761	-7.5	40,400
	1257	1938	712	0.0	42,900
	1258	1806	718	-1.0	
	1250	1727	715	-2.0	42,600
	1260	1629	713	3.0	42,700 42,800
	1261	1555	717	4.0	42,600
	1262	1468	717	·5.0	42,600
	1263	1413	722	-6.0	W. 42:400 See 3
= .	1264	1340	717	-7.0	21.42.600 (2.1.2) All
	1265	1263	717	8.0	42,600
	1266	1182	720	-9.0	42,500
	1267	1110	717	-10.0	42,600
	1268	1055	717	-11.0	42,600
	1269	999	717	-12.0	42,600
	1270	959	715	-13.0	42,700
	1271	905	712	-14.0	42,900
	1272	857	714	-15.0	42,800
	1273	810	705	16.0	43,300
	1274		711	·17:0	42,900
	1277	737	708	-18.0	43,100
	1278	702	711	-19.0	42,900
	1279	671	710	-20.0	43,000
	1280	645	710	-21.0	43,000
	1281 1282	617	707	-22.0	43,100
	1283	595 573	704	-23.0	43,300
	1284	552	700 695	-24.0	43,500
	1285	536	694	-25.0	43,700
	1286	515	687	-26.0 -27.0	43,800 O
	1287	496	683	-27.0	44,200 (). 44,400 ().
	1288	467	669	-29.0	45,200
٠	1289	447	667	-30.9	45,300
. '	1290	427	655	-31.0	45,900
	1291	412	655	-32.0	45,900
	1292	397	652	-33.0	46,100
	1293	381	654	-34.0	46,000
	1294	365	653	-35.0	46,100
•	1295	348	653	<-35.0	46,100

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Table of
Table 2.

POP name	Protein name	WSN.s	Bacic for identification
IDS:3_ALPHA_HDDH	3-a-hydroxysteroid-dihydrodiol- dehydrogenase, an enzyme of	137, 159	ovided by Dr. T.M.
IDS:ACTIN_BETA	steroid metabolism ß cellular actin, a cytoskeletal protein	38	of Madrine, University of Pennsylvania, Homobrane continue with received to the continue with re
IDS:ACTIN_GAMMA	y cellular actin, a cytoskeletal protein	89	Systems Lympolities with second to the secon
IDS:ALBUMIN IDS:APO_A-I	Serum albumin, mature form. Apo A-I plasma lipoprotein, mature form	21, 28, 33 236, 463	Systems Predominance in ret plasma
IDS:CALMODULIN	(fantativa). Calmodulin, an acidic cytosolic calcium-	123, 649	hwering drugs.
IDS:CATALASE	binding protein Catatase (peroxisomal)	54, 61, 106	Presence in nirtied perceived and in the contractions of the contr
IDS:CPKSPOTS	Spots contributed by the CPK charge	1257 - 1295	to mouse catalase
IDS:CPS	Statioards (not rat liver proteins) Carbamoyl phosphate synthase	114, 157, 167, 174, 1184, 1185, 1186, 1222	Pure protein provided by Dr. Margaret Marshall.
IDS:CYTOCHROME_BS	Cytochrame b5	87, 477	Dopartment of Pharmacokogy, Medical School, University of Wisconsin - Madison. Pure potein provided by Dr. Andrew Parkinson, Dopartment of Pharmacology, Toxicology and Therapeutics, University of Kensas Medical
IDS:FABP·L	Liver fatty-acid binding protein	227	Center Pure protein provided by Dr. Nathan Bass, Department of Medicine, University of California School of
IDS:HMG-COA_SYNTHASE	Cytosolic HMG-CoA Symhase	133, 144, 235, 413	Madicine, San Francisco Antibody provided by Dr. Michael Greenspan, Merck Sharp & Dohme Research Laboratories.
IDS:LAMIN_B	Lamin B, a nuclear protein	415, 734	Rahway, NJ Homologous position with respect to other mammallan
IDS:MITCON:1	Mitcon: 1 (F1 ATPase subunit), a	17, 49, 71, 340, 1245, 1246, 1247, 1249	systems Homologous position with respect to other mammalian
IDS:MITCON:2	Mitcon:2, a mitochondrial matrix stress	15, 25, 110, 1241, 1242, 1243, 1244	systems, presence in mitochondria Homologous position with respect to other memmelien
IDS:MITCON:3	Mitcon:3, a mitcohondrial matrix stress	18, 35, 226, 600, 1238, 1239, 1240	systems, presence in mitochondria Homologous position with respect to other mammallan
IDS:NADPH_P450_RED	NADPH cytochrome P-450 reductase, frequently co-induced with P-450's	175, 251, 812	systems, presence in mitochondria Pure protein provided by Dr. Andrew Parkinson, Department of Pharmacology, Toxkoology and Therapeutics, University of Kansas Medical
IDS:PDI	Protein disulphide isomerase 1	168, 1170, 1171, 1172	Center Sequence Information obtained by R.M. Van Frank,
IDS:PLASMA_PROTEINS	Rat plasma proteins observed in liver	21, 28, 33, 44, 72, 102, 115, 197, 236, 246, 248, 257, 293, 332, 347, 364, 369, 419, 432, 463, 468, 518, 552, 605, 623, 666, 667, 725,	Lilly Research Laboratories, Indianapolis Piasma coelectrophoresis studies
IDS:PRO-ALBUMIN	Serum albumin precursor		Relative position to mature albumin, presence in micro-
IDS:PYRCARBOX IDS:SOD	Pyruvate carboxylase Superoxide dismutase	179, 1180, 1181, 1182, 1183 135	somes Pavlica, R.J., et al., BBA (1990) <i>1022</i> 115-125. Sequence Information obtained by R.M. Van Frank,
IDS:TUBULIN_ALPHA	a tubulin, a cytoskeletal protein	56, 132, 1224, 1252	Lily Research Laboratories, Indianapolis Homologous position with respect to other mammalian
IDS:TUBULIN_BETA	B tubulin, a cytoskeletal protein	50, 1225, 1226, 1251	systems Homologous position with respect to other mammatan systems

Table 3. Computed pl's of two sets of carbamylated protein standards: Rabbit muscle CPK and human hemoglobin (Hb)

	nemoglobin (No)										
	Protein Name	PIR Name	#ASP 3.9	#GLU :4.1	#HIS	#LYS	#ARG	NH2- 7.0	Calc	Real	
0	Rabbit muscle CPK	KIRBCM	28	27	17	34					
-1	•		28	27	17	33	18 18	1	6.84 6.67	0.0 -1	
. —	Sold Barriers Te	e et ingres in	. 28		17						n Maria Santa and Maria Maria
	compared to other property			. 27		31	18	· • • • • • • • • • • • • • • • • • • •	:6.42	. , 3	Marie Carlos de la Marie M A marie M
	356 Set 2011, 1875-197.	50%, .		- , 27		30	18	1	6.31	-4	
-5 -6	e de la granda de la companya de la La companya de la co	··· • •	. 28 28	27 27	17 17	29	18	1	6.21	-5	
- 7			28	27	17	28 27	18 18	1	6.12 6.03	-6 -7	
	orra (n. 15 eta 16 il.)	or with great	28	27-	. 17	:: 26	18	€0. 1 /2	i.5.94	/- هـ ،	of the medical process
9	arthur buston is a	di	28	. 27	.17	25	18	. 1.,	5.85	-9	
iu ia 🕬	Branch Barrell		28° 28°	27	17	24	18	1	5.76	-10	(1) (1) (4) (4) (4) (4) (4) (4) (4) (4) (4) (4
	energy of the same		28	: 27 - 27 :	· 17 - 17 :	23		1 2 91 1			menter i sente fili de recomentatione. Als
3 .	MED HARADA MAR	., , , ,	28		17		:.18. :-18.	. 1 . 1	5.48 5.48	P 7-12	e training the property of the year
4 .			28	27	17.	20	18	•	5.39	, -13, -14	one may be with seve
5			28	27	17	19	18	'' i	5.29	-15	order to a live to the state
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8	· to our		28	. 27 27	17 17	17.		- 199		·;•17	or articles of the second
9			28	27	17	16 15	. 18 18	1 1 1 1	5.04 4.96	-18 -19	of the first of the company of the second sections
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1		****	28	. 27	: 17	13	18		4.83	-21	
2 3	·	٠, .	28 28	27	17	12	18	1	4.77	-22	
4 .		1.	28	27 27	17 17	11 10 °	18	1	4.71	-23	the same of the same of the same
5			28	27	17	. 9	18 18	1	4.66 4.61	-24 -25	discovering and then
5	Control of the contro		28	27	17	. 8	18	1	4.56	-25 -26	Commence of the second
7 3			28	27	17	· 7	18	1	4.52	-27	Control of the Contro
,)			28 28	27 27	17	6.		1 :	4.48	-28	· 网络大家 对自己的 4 50
)			28	.27	17 17	5 4	18 18		4.44	-29	A Commence of the Commence of the
ì .			28	27	17	3	18	1, 1	4.40	-30 -31	
?		'	28	27	17	٠ 2	_	-	4.32	-31 -32	T
) - 2 - I	and the second		28	27	.17	. 1	18	14	4.29	-33	
↓, 5	****		28 28	27.	17	. 0	18		4.25	-34	
<u> </u>	* * * * * * * * * * * * * * * * * * * *	<u></u>		27	17	0	18	0	4.22	-35	The second secon
) , , (Hb-beta, human 🦠 🖠	HBHU	~7 · <u>;</u>	8	9	11 5	. 3	1	7.18		Ar and
		• • •	7.	. 8	9	10	3	1	6.79		
٠.			.7 7	8	9	9	3	1	6.53	-1.8	
			7	8	9	8 7	3 3	1	6.32	-3.2	• • • • • • • • • • • • • • • • • • • •
i., .	•		.7	8	9	6	3	1	6.13 5.96	-5.3 -7.2	
i .			7	8	9	5	3	1	5.78	-7.2	*
			7	8	9	4	3	1 .	5.59	-12.3	
•	•		7	8	9	3	3	1	5.37	-15.5	
			7 7	8 8	9	2	3	1	5.14	-18.0	•
	•		7	8	9 9	1	3	1	4.91	-21.0	•
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Table 4. Computed prs of some known proteins related to measured CPK prs

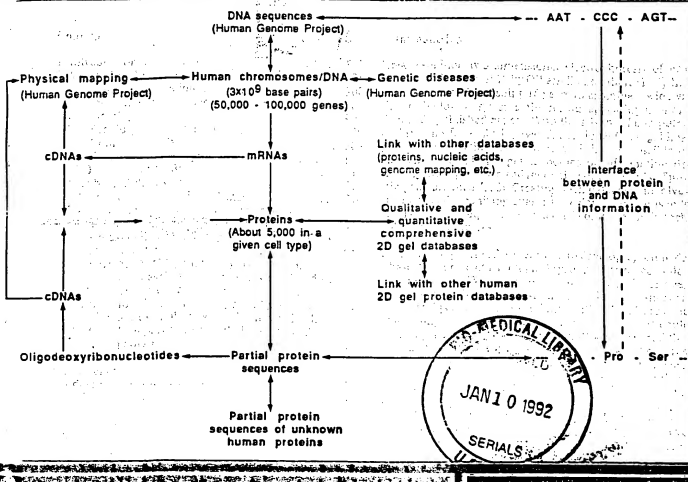
1 - 42 (1)

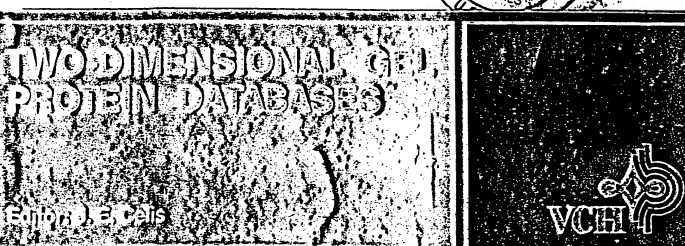
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Protein Name	PIR Name	#ASP 3.9	#GLU 4.1	#HIS 6.0		#ARG 12.5	Calc pl	Real
0.44 A Creatine phospho kinase (CPK), rabbit muscle	KIRBCM	28,	27.	17	34	18	6.84	0.0
1 Fatty acid-binding protein, rat hepatic	FZRTL	., , 5 .,		~2	. 16	2	7.83	-3.0
2 b2-microglobulin, human	MGHUB2	7	8	4	8	5	6.09	-5.0
3 Carbamoyl-phosphate synthase, rat	SYRTCA	72	96'	28	95	56	5.97	-5.5
4 Proalbumin (serum albumin precursor), rat	ABRTS	32	57	· 15.	⊹: 53	27 .6	5.98	6.2
5 Serum albumin, rat	ABRTS	32	5 7	. 15	· 53	. 24	5.71	-9.0
6 Superoxid dismutase (Cu-Zn, SOD), rat	A26810	8	11	10	9	4	5.91	-9.2
7 Phospholipase C, phophoinositide-specific (?), rat	A28807	34	42	9	49	21	5.92	-9.2
8 Albumin, human	ABHUS "	36	61	16	60	24	5.70	-11.9
g Apo A-I lipoprotein, rat	A24700	18	24	6	·c 23:	12	· 5.32	≈:±13.7
10 proApo A-I lipoprotein, human	LPHUA1	16	30	:6	2,1	17	5.35	-14.3
NADPH cytochrome P-450 reductase, rat	RDRTO4	41:	60.	21	.38	. 36	5.07	-15.6
12 Retinol binding protein, human	VAHU	18	10	2	10	14	5.04	-16.9
13 Actin beta, rat	ATRTC	23	26	9	19	" 18	5.06	-17.2
14 Actin gamma, rat	ATRTC	20	29	• • • • • •	· · · · 19	18	5.07	-16.8
15 Apo A-l'lipoprotein, human	LPHUA1	··· '16"	30	5	- 21	: .16	··· 5.10	-17.5
16 Apo A-IV lipoprotein, human	LPHUA4	20	49.	8	28	- 24	4.88	-19.7
17 Tubulin alpha, rat	UBRTA	27	37	13	19	21	4.66	-19.8
18 F1ATPase beta, bovine	PWBOB	25	36	9	22	22	4.80	-21.0
19 Tubulin beta, pig	UBPGB	26	36	10	15	22	4.49	-22.5
20 Protein disulphide isomerase (PDI), rat hepatic	ISRTSS	43	51	11	51	9	4.07	-25.0
21 Cytochrome b5, rat	CBRT5	55 10	15	6.	10	4	4.59	-26.0
22 Apo C-II lipoprotein, human	LPHUC2	: 4	7	. 0	6	1:	4.44	-30.5
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Amino acid pl assumed in calulation:		3.9	4.1	6.0	10.8	12.5	1 1 2 4t	1.1

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N. Leigh Anderson Ricardo Esquer-Blasco. Jean-Paul Holmann Norman, G. Anderson

A two-dimensional gel database of rat liver proteins useful in gene regulation and drug effects studies

Large Scale Biology Corporation, Rockville, MD

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Abbreviations: CBB, Coomassie Brilliant Biue; CPK, creatine phosphokinase; 2-D, two-dimensional; IEF, isoelectric focusing; MSN, master spot number; NP-40, Nonidet P-40, SDS, sodium dodecyl sulfate

1 Introduction is a compact of a manager for the compact of the co

High-resolution two-dimensional electrophoresis of proteins, introduced in 1975 by O'Farrell and others [1-4], has been used over the ensuing 16 years to examine a wide variety of biological systems, the results appearing in more than 5000 published papers. With the advent of computerized systems for analyzing two-dimensional (2-D) gel images and constructing spot databases, it is also possible to plan and assemble integrated bodies of information describing the appearance and regulation of thousands of protein gene products [5, 6]. Creating such databases involves amassing and organizing quantitative data from thousands of 2-D gels, and requires a substantial commitment in technology and resources.

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Given the long-term effort required to develop a protein database, the choice of a biological system takes on considerable importance. While in vitro systems are ideal for answering many experimental questions, especially in cancer research and genetics, our experience with cell cultures and tissue samples suggests that some in vivo approaches could have major advantages. In particular, we have noticed that liver tissue samples from rats and mice appear to show greater quantitative reproducibility (in terms of individual protein expression) than replicate cell cultures. This is perhaps a natural result of the homeostasis maintained in a complete animal vs. the well-known variability of cell cultures, the latter due principally to differences in reagents (e.g., fetal bovine serum), conditions (e.g., pH) and genetic "evolution" of cell lines while in culture. It is also more difficult to generate adequate amounts of protein from cell culture systems (particularly with attached cells), forcing the investigator to resort to radioisotope-based or silver-based staindetection methods. While these methods are more sensitive (sometimes much more sensitive) than the Coomassie Brilliant Blue (CBB) stain typically used for protein detection in "large" protein samples, they are generally more variable, more labor-intensive and, in the case of radiographic methods, may generate highly "noisy" images, due to the properties of the films used. By contrast, large protein samples can easily be prepared from liver using urea/Nonidet P-40 (NP-40) solubilization and stained with CBB, which has the advantage of being easily reproducible [8]. Finally, there remains the question of the "truthfulness" of many in vitro systems as compared to their in vivo analogs; how great are the changes caused by the introduction into a culture and the associated shift to strong selection for growth, and how do these affect experimental outcomes? Hence the apparent advantages of in vitro systems, in terms of experimental manipulation, may be counterbalanced by other factors relating to 2-D data quality.

There is a second important class of reasons for exploring the use of an in vivo biological system such as the liver. Historically, there have been two broad approaches to the mechanistic dissection of biochemical processes in intact cellular systems: genetics (a search for informative mutants) and the use of chemical agents (drugs and chemical toxins). Both approaches help us to understand complex systems by disrupting some specific functional element and showing us the result. With the development of techniques for genetic manipulation and cloning, the genetic approach can be effectively applied either in vitro or in vivo, although the in vitro route is usually quicker. The chemical approach. can also be applied to either sort of biological system; here, however, the bulk of consistently acquired information is in experimental animals (rats and mice). While most biologists know a short list of compounds having specific, expenmentally useful effects (e.g., inhibitors of protein synthesis, ionophores, polymerase inhibitors, channel blockers, nucleotide analogs, and compounds affecting polymerization of cytoskeletal proteins), there is a much larger number of interesting chemically-induced effects, most of them characterized by toxicologists and pharmacologists in rodent systems. Just as a thorough genetic analysis would involve saturating a genome with mutations, it is possible to imagine a saturating number of drugs, the analysis of whose actions would reveal the complete biochemistry of the cell. While organized drug discovery efforts usually target specific desired effects, the nature of the process, with its dependence on screening large numbers of compounds, necessarily produces many unanticipated effects. It is therefore reasonable to suppose that the required broad range of compounds necessary to achieve "biochemical saturation" may be forthcoming; in fact, it may already exist among the hundreds of thousands of compounds that failed to qualify as drugs.

Among organs, the liver is an obvious choice for the study of chemical effects because of its well-known plasticity and responsiveness. The brain appears to be quite plastic (e.g. [7]), but it is a complicated mixture of cell types requiring skillful dissection for most experiments. The kidney, while quite responsive, also presents a potentially confounding mixture of cell types. The liver, by contrast, is made up of one predominant cell type which is easy to solubilize; the hepatocyte, representing more than 95% of its mass. Most importantly, the liver performs many homeostatic functions that require rapid modulation of gene expression. It appears that most chemical agents tested affect gene expression in the liver at some dosage (N. Leigh Anderson, unpublished observations), an interesting contrast to our earlier work with lymphocytes, for example, which seem to be much less responsive. Such results conform to the expectation that cells with a homeostatic, physiological role should be more plastic than cells differentiated for a purpose dependent on the action of a limited number of specific genes.

The liver also allows the parallels between in vitro and in vivo systems to be examined in detail. Significant progress

has been made in the development of mouse, rat and human hepatocyte culture systems, as well as in precision-cut tissue slices. Using such an array of techniques, it is possible to assemble a matrix of mammalian systems including mouse and rat in vivo on one level and mouse, rat and human in vitro on a second level, and to compare effects between species and between systems. This approach allows us to draw informed conclusions regarding the biochemical universality of biological responses among the mammals, and to offer some insight into the validity of in vitro approaches for toxicological screening. We believe this data will be necessary if in vitro alternatives are to achieve wide usage in government-mandated safety testing of drugs, consumer products and industrial and agricultural chemicals.

A number of interesting studies have been published using 2-D mapping to examine effects in the rodent liver. A number of investigarors have made use of the technique to screen for existing genetic variants [8-11] or induced mutations [12-14], mainly in the mouse. This work builds on the wealth of genetic information available on the mouse and its established position as a mammalian mutation-detection system. While some studies of chemical effects have been undertaken in the mouse [15-17], most have used the rat [18-23]. The examination of the cytochrome p-450 system, in particular, has been carried out almost exclusively on the rat [24, 25].

These considerations lead us to conclude that rodent liver offers the best opportunity to systematically examine an array of gene regulation systems, and ultimately to build a predictive model of large-scale mammalian gene control. The basic underlying foundation of such a project is a reliable, reproducible master 2-D pattern of liver, to which ongoing experimental results can be referred. In this paper, we report such a master pattern for the acidic and neutral proteins of rat liver (pattern F344MST3). In future, this master will be supplemented by maps of basic proteins, and analogous maps of mouse and human liver.

2 Materials and methods

2.1 Sample preparation

Liver is an ideal sample material for most biochemical studies, including 2-D analysis. A sample is taken of approximately 0.5 g of tissue from the apical end of the left lobe of the liver. Solubilization is effected as rapidly as practical; a delay of 5-15 min appears to cause no major alteration in liver protein composition if the liver pieces are kept cold (e.g., on ice) in the interim. In the solubilization process, the liver sample is weighed, placed in a glass homogenizer (e.g., 15 mL Wheaton); 8 volumes of solubilizing solution.

The solubilizing solution is composed of 2 % NP-40 (Sigma), 9 m urea (analytical grade, e.g., BDH or Bio-Rad), 0.5% dithiothreitol (DTT; Sigma) and 2% carrier ampholytes (pH 9-11 LKB: these come as a 20% stock solution, so 2% final concentration is achieved by making the final solution 10% 9-11 Ampholine by volume). A large batch of solubilizer (several hundred mL) is made and stored frozen at -80°C in aliquots sufficient to provide enough for one day's estimated sample preparation requirement. The solution is never allowed to become warmer than room temperature at any stage during preparation or thawing for use, since heating of concentrated urea solutions can produce contaminants that covalently modify proteins producing artifactual charge shifts. Once thawed, any unused solubilizer is discarded.

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is added (i.e., 4 mL per 0.5 g tissue) and the mixture is homogenized using first, the loose- and then then the tight-fitting glass pestle. This takes approximately 5 strokes with each pestle and is carried out at room temperature because urea would crystallize out in the cold. Once the liver sample is thoroughly homogenized in the solubilizer, it is assumed that all the proteins are denatured (by the chaotropic effect of the urea and NP-40 detergent) and the enzymes inactivated by the high pH (-9.5). Therefore these samples may be kept at room temperature until they can be centrifuged or frozen as a group (within several hours of preparation). The samples are centrifuged for 6×10^6 g min (e.g., 500 000 × g for 12 min using a Beckman TL-100 centrifuge). The centrifuge rotor is maintained at just below room temperature (e.g., 15-20°C), but not too cold, so as to prevent the precipitation of urea. The centrifuge of choice is a Beckman TL-100 because of the sample tube sizes available, but any ultracentrifuge accepting smallish tubes will suffice. When: an appropriate centrifuge is not available near the site of sample preparation, samples can be frozen at -80°C and thawed prior to centrifugation and collection of supernatants. Each supernatant is carefully removed following centrifugation and aliquoted into at least 4 clean tubes for storage. This is done by transferring all the supernatant to one clean tube, mixing this gently (to assure homogeneous composition) and then dividing it into 4 aliquots. The aliquots are frozen immediately at -80°C. These multiple aliquots can provide insurance against a failed run or a freezer breakdown.

2.2 Two-dimensional electrophoresis

Sample proteins are resolved by 2-D electrophoresis using the 20×25 cm lso-Dalt* 2-D gel system ([26-29]; produced by LSB and by Hoefer Scientific Instruments, San Francisco) operating with 20 gels per batch. All first-dimensional isoelectric focusing (IEF) gels are prepared using the same single standardized batch of carrier ampholytes (BDH 4-8A in the present case, selected by LSB's batchtesting program for rat and mouse database work**). A 10 µL sample of solubilized liver protein is applied to each gel, and the gels are run for 33 000 to 34 500 volt-hours using a progressively increasing voltage protocol implemented by a programmable high-voltage power supply. An Angelique" computer-controlled gradient-casting system (produced by LSB) is used to prepare second-dimensional sodium dodecyl sulfate (SDS) polyacrylamide gradient slab gels in which the top 5% of the gel is 11%T acrylamide, and the lower 95% of the gel varies linearly from 11% to 18%T.

This system has recently been modified so as to employ a commercially available 30.8%T acrylamide/N,N-methylenebisacrylamide prepared solution (thus avoiding the handling of the solid acrylamide monomer) and three additional stock solutions: buffer (made from Sigma pre-set Tris), persulfate and N,N,N,N-tetramethylethylenediamine (TEMED). Each gel is identified by a computer-printed filter paper label polymerized into the lower left corner of the gel. First-dimensional IEF tube gels are loaded

directly (as extruded) onto the slab gels without equilibration, and held in place by polyester fabric wedges (Wedgies", produced by LSB) to avoid the use of hot agarose. Second-dimensional slab gels are run overnight, in groups of 20, in cooled DALT tanks (10°C) with buffer circulation. All run parameters, reagent source and lot information, and notations of deviation from expected results are entered by the technician responsible on a detailed, multi-page record of the experiment.

2.3 Staining

Following SDS-electrophoresis, slab gels are stained for protein using a colloidal Coomassie Blue G-250 procedure in covered plastic boxes, with 10 gels (totalling approximasely I L of gel) per box. This procedure (based on the work of Neuhoff [30, 31]) involves fixation in 1.5 L of 50% ethanol and 2% phosphoric acid for 2 h, three 30 min washes, each in 2L of cold tap water, and transfer to 1.5 L of 34% methanol, 17% ammonium sulfate and 2% phosphoric acid for 1 h, followed by the addition of a gram of powdered Coomassie Blue G-250 stain. Staining requires approximately 4 days to reach equilibrium intensity, whereupon gels are transferred to cool tap water and their surfaces rinsed to remove any particulate stain prior to scanning. Gels may be kept for several months in water with added sodium azide. The water washes remove ethanol that would dissolve the stain (and render the system noncolloidal, with high backgrounds). The concentrated ammonium sulfate and methanol solution is diluted by equilibration with the water volume of the gels to automatically achieve the correct final concentrations for colloidal staining. Practical advantages of this staining approach can be summarized as follows: (i) the low, flat background makes computer evaluation of small spots (max OD < 0.02) possible, especially when using laser densitometry; (ii) up to 1500 spots can be reliably detected on many gels (e.g., rat liver) at loadings low enough to preserve excellent resolution; and (iii) reproducibility appears to be very good: at least several hundred spots have coefficients of reproducibility less than 15%. This value is at least as good as previous CBB methods, and significantly better than many silver stain systems.

2.4 Positional standardization

The carbamylated rabbit muscle creatine phosphokinase (CPK) standards [32] are purchased from Pharmacia and BDH. Amino acid compositions, and numbers of residues present in proteins used for internal standardization, are taken from the Protein Identification Resource (PIR) sequence database [33].

2.5 Computer analysis

Stained slab gels are digitized in red light at 134 micron resolution, using either a Molecular Dynamics laser scanner (with pixel sampling) or an Eikonix 78/99 CCD scanner. Raw digitized gel images are archived on high-density DAT tape (or equivalent storage media) and a greyscale videoprint prepared from the raw digital image as hard-copy backup of the gel image. Gels are processed using the Kepler software system (produced by LSB), a commercially available workstation-based software package built on

This material (succeeding certified batches of which are available from Hoefer Scientific Instruments) has the most linear pH gradient produced by any ampholyte tested except for the Pharmacia wide range (which has an unacceptable tendency to bind high-molecular weight acidic proteins, causing them to streak).

some of the principles of the earlier TYCHO system [34–41]. Procedure PROC008 is used to yield a spotlist giving position, shape and density information for each detected spot. This procedure makes use of digital filtering, mathematical morphology techniques and digital masking to remove the background, and uses full 2-D least-squares optimization to refine the parameters of a 2-D Gaussian shape for each spot. Processing parameters and file locations are stored in a relational database, while various log files detailing operation of the automatic analysis software are archived with the reduced data. The computed resolution and level of Gaussian convergence of each gel are inspected and archived for quality control purposes.

Experiment packages are constructed using the Kepler experiment definition database to assemble groups of 2-D patterns corresponding to the experimental groups (e.g., treated and control animals). Each 2-D pattern is matched to the appropriate "master" 2-D pattern (pattern F344MST3 in the case of Fischer 344 rat liver), thereby providing linkage to the existing rodent protein 2-D databases. The software allows experiments containing hundreds of gels to be constructed and analyzed as a unit, with up to 100 gels displayed on the screen at one time for comparative purposes and multiple pages to accommodate experiments of > 1000 gels. For each treatment, proteins showing significant quantitative differences vs. appropriate controls are selected using group-wise statistical parameters (e.g., Student's t-test, Kepler procedure STUDENT). Proteins satisfying various quantitative criteria (such as P< 0.001 difference from appropriate controls) are represented as highlighted spots onscreen or on computer-plotted protein maps and stored as spot populations (i.e., logical vectors) in a liver protein database. Quantitative data (spot parameters, statistical or other computed values) are stored as real-valued vectors in the database. Analysis of coregulation is performed using a Pierson product-moment correlation (Kepler procedure CORREL) to determine whether groups of proteins are coordinately regulated by any of the treatments. Such groups can be presented graphically on a protein map, and reported together with the statistical criteria used to assess the level of coregulation. Multivariate statistical analysis (e.g., principal components' analysis) is performed on data exported to SAS (SAS Institute).

2.6 Graphical data output

Graphical results are prepared in GKS and translated within Kepler® into output for any of a variety of devices. Linedrawing output is typically prepared as Postscript and printed on an Apple Laserwriter. Detailed maps presented here have been generated using an ultra-high-resolution Postscript-compatible Linotronic output device. Greyscale graphics are reproduced from the workstation screen using a Seikosha videoprinter. Patterns are shown in the standard orientation, with high molecular mass at the top and acidic proteins to the left.

2.7 Experiment LSBC04

In the study described here 12-week-old Charles River male F344 rats were used. Diets were prepared at LSB, based on a Purina 5755M Basal Purified Diet. Lovastatin and cholestyramine were obtained as prescription pharma-

ceuticals, ground and mixed with the diet at concentrations. of 0.075% and 1%, respectively. The high cholesterol diet was Purina 5801M-A (5% cholesterol plus 1% sodium cholate in the control diet). Animal work was carried out by Microbiological Associates (Bethesda, MD). Animals were acclimatized for one week on the control diet, fed test or control diets for one week, and sacrificed on day 8. Average daily doses of lovastatin and cholestyramine in appropriate. groups were 37 mg/kg/day and 5 g/kg/day, respectively, based on the weight of the food consumed. Liver samples were collected and prepared for 2-D electrophoresis according to the standard liver protocol (homogenization in 8 volumes of 9 m urea, 2% NP-40, 0.5% dithiothreitol, 2% LKB pH 9-11 carrier ampholytes, followed by centrifugation for 30 min at 80 000 × g). Kidney, brain and plasma. samples were frozen. Gels were run as described above, and the data was analyzed using the Kepler system. Gels were scaled, to remove the effect of differences in protein loading, by setting the summed abundances of a large number of matched spots equal for each gel (linear scaling).

3 Results and discussion-

3.1 The rat liver protein 2-D map

- F344MST3 is a standard 2-D pattern of rat liver proteins, based on the Fischer 344 strain. This pattern was initiated from a single 2-D gel and extensively edited in an experiment comparing it to a range of protein loads, so as to include both small spots and well-resolved representations of high-abundance spots. More than 700 rat liver 2-D patterns have been matched to F344MST3 in a series of drug effects and protein characterization experiments, and numerous new spots (induced by specific drugs, for instance) have been added as a result. A modified version including additional spots present in the Sprague-Dawley outbred rat has also been developed (data not shown). Figure 1 shows a greyscale representation and Fig. 2 a schematic plot of the master pattern. More than 1200 spots are included, most of which are visible on typical gels loaded with 10 µL of solubilized liver protein prepared by the standard method and stained with colloidal Coomassie Blue. Master spot numbers (MSN's) have been assigned to all proteins, and appear in the following figures, each showing one quadrant of the pattern. Figure 3 shows the upper left (acidic, high molecular mass) quadrant, Fig. 4 the upper right (basic, high molecular mass) quadrant, Fig. 5 the lower left (acidic, low molecular mass) quadrant, and Fig. 6 the lower right (basic, low molecular mass) quadrant. The quadrants overlap as an aid to moving between them. The gel position (in 100 micron units), isoelectric point (relative to the CPK internal pl standards) and SDS molecular mass (from the calibration curve in Fig. 8) are listed for each spot (Table 1). Because of the precision of the CPK-pl values, these parameters can be used to relate spot locations between gel systems more reliably than using pl measurements expressed as pH. A major objective of current studies is the identification of all major spots corresponding to known liver proteins, as well as rigorous definitions of subcellular organelle contents. Of particular interest to us is the parallel development of identifications in the rat and mouse liver maps, allowing detailed comparisons of gene expression effects in the two systems. The results of these studies will be presented systematically in a later edition of this database,

but we include here a useful series of 22 orienting identifications as an aid to other users of the rat liver pattern (Table 2).

3.2 Carbamylated charge standards, computed p/s and molecular mass standardization

We have previously shown that the use of a system of close-ly-spaced internal pl markers (made by carbamylating a basic protein) offers an accurate and workable solution to the problem of assigning positions in the pl dimension [32]. The same system, based on 36 protein species made by carbamylating rabbit muscle CPK, has been used here to assign pl's to most rat liver acidic and neutral proteins. The standards were coelectrophoresed with total liver proteins, and the standard spots added to a special version of the master pattern F344MST3. The gel X-coordinates of all liver protein spots lying within the CPK-charge train were then transformed into CPK pl positions by interpolation between the positions of immediately adjacent standards (Table 1) using a Kepler vector procedure.

It has proven possible to compute fairly accurate p/values for many proteins from the amino acid composition [42]. We have attempted here to test a further elaboration of this approach, in which we computed prs for the CPK standards themselves, based on our knowledge of the rabbit muscle CPK sequence and the fact that adjacent members of the charge train typically differ by blockage of one additional lysine residue (Table 3). We compared these values to similar computed pl's for an additional set of carbamylated standards made from human hemoglobin beta chains and a series of rat liver and human plasma proteins of known position and sequence (Fig. 7, Table 4). The result demonstrates good concordance between these systems. Two proteins show significant deviations: liver fatty-acid binding protein (FABP; #1 in Table 4) and protein disulphide isomerase (#20 in the table). The FABP spot present on F344MST3 may represent a charge-modified version of a more basic parent spot closer to the expected pl, not resolved in the IEF/SDS gel. Of particular importance is the fact that, by comparing computed pl's of sequenced but unlocated proteins with the CPK p/s, we can assign a probable gel location without making any assumptions regarding the actual gel pH gradient. This offers a useful shortcut, given the vagaries of pH measurement on small diameter IEF gels. We have used this approach to compute the CPK prs of all rat and mouse proteins in the PIR sequence database, as an aid to protein identification (data not shown).

In order to standardize SDS molecular weight (SDS-MW), we have used a standard curve fitted to a series of identified proteins (Fig. 8). Rather than using molecular mass per se, we have elected to use the number of amino acids in the polypeptide chain, as perhaps a better indication of the length of the SDS-coated rod that is sieved by the second dimension slab. The resulting values were multiplied by 112 (the weighted average mass of amino acids in sequenced proteins) to give predicted molecular masses. Because we use gradient slabs, we have not constrained the fitted curve to conform to any predetermined model; rather we tried many equations and selected the best using the program "Tablecurve" on a PC. The equation chosen was y = a + bx + c/x', where y is the number of residues, x is the gel

Y coordinate, a is 511.83, b is -0.2731 and c is 33183801. The resulting fit appears to be fairly good over a broad range of molecular mass.

3.3 An example of rat liver gene regulation: Cholesterol metabolism

Experiment LSBC04 was designed as a small-scale test of the regulation of cholesterol metabolism in vivo by three agents included in the diet: lovastatin (Mevacora, an inhibitor of HMG-CoA reductase); cholessyramine (a bile acid sequestrant that has the effect of removing cholesterol from the gut-liver recirculation); and cholesterol itself. The first two agents should lower available cholesterol and the third should raise it, allowing manipulation of relevant gene expression control systems in both directions. Such an experiment offers an interesting test of the 2-D mapping system-since most of the pathway enzymes are present in low abundance, many are membrane-bound and difficult. ito solubilize, and the pathway itself is complex. Approximately 1000 proteins were separated and detected in liver homogenates. Twenty-one proteins were found to be affected by at least one treatment, and these could be divided into several coregulated groups.

3.3.1 MSN 413 (putative cytosolic HMG-CoA synthase) and sets of spots regulated coordinately or inversely

One group of spots (including a spot assigned to the cytosolic HMG-CoA synthase, MSN 413) showed the expected increase in abundance with lovastatin or cholestyramine, the synergistic further increase with lovastatin and cholestyramine, and a dramatic decrease with the high cholesterol diet. Spot number 413 is the most strongly regulated protein in the present experiment, showing a 5- to 10-fold induction after a 1 week treatment with 0.075 % lovastatin and 1% cholestyramine in the diet (Figs. 9 and 10). Its expression follows precisely the expectation for an enzyme whose abundance is controlled by the cholesterol level; it is progressively increased from the control levels by cholestyramine, lovastatin and lovastatin plus cholestyramine, and it sinks below the threshold of detection in animals fed the high cholesterol diet. This spot has been tentatively identified as the cytosolic HMG-CoA synthase, based on a reaction with an antiserum to that protein provided by Dr. Michael Greenspan at Merck Sharp & Dohme Research Laboratories. This enzyme lies immediately before HMG-CoA reductase in the liver cholesterol biosynthesis pathway, and is known to be co-regulated with it. Spot 413 has an SDS molecular weight of about 54 000 and a CPK p/ of -11.4, in reasonably close agreement with a molecular weight of 57300 and a CPK pl of -15.7 computed from the known sequence of the hamster enzyme [43].

Using a classical product-moment correlation test (Kepler procedure CORREL), a series of five additional spots was found to be coregulated with 413. The level of correlation was exceedingly high (> 95%). Two of these, 1250 and 933, are at similar molecular weights and approximately one charge more acidic than 413 (Fig. 9), indicating that they may be covalently modified forms of the 413 polypeptide. This suspicion is strengthened by the observation that both spots are also stained by the antibody to cytosolic HMG-CoA synthase. The remaining three correlated spots appear

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to comprise an additional related pair (1253 and 1001) of around 40 kDa and a single spot (1119) of around 28 kDa. Because these two presumed proteins are present at substantially lower abundances than 413, and because the cytosolic HMG-CoA synthase is reponed to consist of only one type of polypeptide, they are likely to represent other, very lightly coregulated enzymes. A second group of six spots was selected based on a regulatory pattern close to the inverse of that for spot 413 (MSN's 34, 79, 178, 182, 204, 347; data not shown). For these proteins, the lowest-level of expression occurs with exposure to lovastatin plus cholestyramine and the highest level upon exposure to the high-cholesterol diet. Spots 182 and 79 are highly correlated and lie about one charge apart at the same molecular weight; they may thus be isoforms of a single protein. The other four spots probably represent additional enzymes or subunits.

3.3.2 MSN 235 and coregulated spots

A third group of five spots, mainly comprised of mitochondrial proteins including putative mitochondrial HMG-CoA synthase spots, showed a modest induction by lovastatin alone, but little or no effect with any of the other treatments (including the combination of lovastatin and cholestyramine; Fig. 12). This result is intriguing because lovastatin was expected to affect only the regulation of enzymes of cholesterol synthesis, which is entirely extra-mitochondrial. Three of the spots (235, 134, 144) form a closelypacked triad at approximately 30 kDa, and are likely to represent isoforms of one protein. All three spots are stained by an antibody to the mitochondrial form of HMG-CoA synthase obtained from Dr. Greenspan. Subcellular fractionation indicates a mitochondrial location. The other two spots (633 at about 38 kDa and 724 at about 69 kDa) are each present at lower abundance than the members of the

3.3.3 An example of an anti-synergistic effect

A sixth spot (367) shows strong induction by lovastatin (two- to threefold), and about half as much induction with lovastatin plus cholestyramine, but without sharing the animal-animal heterogeneity pattern of the 235-set (Fig. 13). This protein is also mitochondrial, and represents the clearest example of an anti-synergistic effect of lovastatin and cholestyramine. The existence of such an effect demonstrates that lovastatin and cholestyramine do not act exclusively through the same regulatory pathway.

3.3.4 Complexity of the cholesterol synthesis pathway

Taken together, these results suggest that treatment with lovastatin alone can affect both cytosolic and mitochondrial pathways using HMG-CoA, while cholestyramine, on the other hand, either alone or in combination with lovastatin, produces a strong effect on the putative cytosolic pathway, but little or no effect on the putative mitochondrial pathway. An explanation for this difference may lie in lovastatin's effect on levels of HMG-CoA and related precursor compounds that are exchanged between the cytosol and the mitochondrion, whereas cholestyramine should affect only the cytosolic pathways directly controlled by cholesterol and bile acid levels. It remains to be explained why some

proteins of the putative mitochondrial pathway are so much more variable in their expression in all groups. An examination of all the coregulated groups suggests that quantitative statistical techniques can extract a wealth of interesting information from large sets of reproducible gels. The abundance of spots in the 413 coregulation group, for example, shows an amazing level of concordance in their relative expression among the five individuals of the lovastatin and cholestyramine treatment group. This effect is not due to differences in total protein loading, since they have already been removed by scaling, and since proteins with quite different regulation patterns can be demonstrated (e.g., Fig. 13). Such effects raise the possibility that many gene coregulation sets may be revealed through the study of a sufficiently large population of control animals (i.e., without any experimental manipulation). This approach, exploiting natural biological variation in protein expression instead of drug effects, offers an important incentive for the construction of a large library of control animal patterns.

4 Conclusions

Because of the widespread use of rat liver in both basic biochemistry and in toxicology, there is a long-term need for a comprehensive database of liver proteins. The rat liver master pattern presented here has proven to be an accurate representation of this system, having been matched to more than 700 gels to date. As the number of proteins identified and the number of compounds tested for gene expression effects grows, we expect this database to contribute valuable insights into gene regulation. Its practical utility in several areas of mechanistic toxicology is already being demonstrated.

Received September 11, 1991

5 References

- [1] O'Farrell, P., J. Biol. Chem. 1975, 250, 4007-4021.
- [2] Klose, J., Humangenetik 1975, 26, 231-243.
- [3] Scheele, G. A., J. Biol. Chem. 1975, 250, 5375-5385.
- [4] Iborra, G. and Buhler, J. M., Anal. Biochem. 1976, 74, 503-511.
- [5] Anderson, N. G. and Anderson, N. L., Behring. Inst. Mitt. 1979, 63, 169-210.

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- [6] Anderson, N. G. and Anderson, N. L., Clin. Chem. 1982, 28, 739-748.
- [7] Heydorn, W. E., Creed, G. J. and Jacobowitz, D. M., J. Pharmacol. Exp. Therap. 1984, 229, 622-628.
- [8] Anderson, N. L., Nance, S. L., Tollaksen, S. L., Giere, F. A. and Anderson, N. G., Electrophoresis 1985, 6, 592-599.
- 191 Racine, R. R. and Langley, C. H., Biochem. Genet. 1980, 18, 185-197.
- [10] Klose, J., Mol. Evol. 1982, 18, 315-328.
- [11] Neel, J. V., Baier, L., Hanash, S. and Erickson, R. P., J. Hered. 1985, 76, 314-320.
- [12] Marshall, R. R., Raj, A. S., Grant, F. J. and Heddle, J. A., Can. J. Genet. Cytol. 1983, 25, 457-446.
- [13] Taylor, J., Anderson, N. L., Anderson, N. G., Gemmell, A., Giometti, C. S., Nance, S. L. and Tollaksen, S. L., in: Dunn, M. J. (Ed.), Electrophoresis '86, Verlag Chemie, Weinheim 1986, pp. 583-587.
- [14] Giometti, C. S., Gemmell, M. A., Nance, S. L., Tollaksen, S. L. and Taylor, J., J. Biol. Chem. 1987, 262, 12764-12767.
- [15] Anderson, N. L., Giere, F. A., Nance, S. L., Gemmell, M. A., Tollaksen, S. L. and Anderson, N. G., in: Galteau, M.-M. and Siest, G. (Eds.), Progrès Récents en Electrophorèse Bidimensionelle, Presses Universitaires de Nancy, Nancy 1986, pp. 253-260.
- [16] Anderson, N. L., Swanson, M., Giere, F. A., Tollaksen, S., Gemmell, A., Nance, S. L. and Anderson, N. G., Electrophoresis 1986, 7, 44–48.

- [17] Anderson, N. L., Giere, F. A., Nance, S. L., Gemmell, M. A., Tollaksen, S. L. and Anderson, N. G., Fundam, Appl. Toxicol. 1987, 8, 39-50.
- [18] Anderson, N. L., in: New Horizons in Toxicology, Eli Lilly Symposium, 1991, in press.
- [19] Antoine, B., Rahimi-Pour, A., Siest, G., Magdalou, J. and Galteau, M. M., Cell. Biochem. Funct. 1987, 5, 217-231.
- [20] Elliott, B. M., Ramasamy, R., Stonard, M. D. and Spragg, S. P. Biochim. Biophys. Acta 1986, 870, 135-140.
- [21] Huber, B. E., Heilman, C. A., Wirth, P. J., Miller, M. J. and Thorgeirs's son, S. S., Hepatology 1986, 6, 209-219.
- . [22] Wirth, P. J. and Vesterberg, O., Electrophoresis 1988, 9, 47-53
- [23] Witzmann, F. A. and Parker, D. N., Toxicol. Lett. 1991, 57, 29-36.
- [24] Rampersaud, A., Waxman, D. J., Ryan, D. E., Levin, W. and Walz, F. G., Jr., Arch. Biochem. Biophys. 1985, 243, 174-183.
- [25] Vlasuk, G. P. and Walz, F. G., Jr., Anal. Biochem. 1980, 105, 112-120.
- [26] Anderson, N. G. and Anderson, N. L., Anal. Biochem. 1978, 85, 351-
- [27] Anderson, N. L. and Anderson, N. G., Anal. Biochem. 1978, 85, 341—354.
- [28] Anderson, L., Hofmann, J.-P., Anderson, E., Walker, B. and Anderson, N. G., in: Endler, A. T. and Hanash, S. (Eds.), Two-Dimensional Electrophoresis, VCH Verlagsgesellschaft, Weinheim 1989, pp. 288-297.
- [29] Anderson, L., Two-Dimensional Electrophoresis: Operation of the ISO-DALT[®] System. Large Scale Biology Press, Washington, DC 1988, ISBN 0-945532-00-8, 170pp.
- [30] Neuhoff, V., Stamm, R. and Eibl, H., Electrophoresis 1985, 6, 427-448.

- [31] Neuhoff, V., Arold, N., Taube, D. and Ehrhardt, W., Electrophoresis 1988, 9, 255-262.
- [32] Anderson; N. L. and Hickman, B.J., Anal. Biochem. 1979, 93, 312-320.
- [33] Sidman, K. E., George, D. E., Barker, W. C. and Hunt, L. T., Nucl. Acids Res. 1988, 16, 1869—1871.
- [34] Taylor, J., Anderson, N. L., Coulter, E. P., Scandora, A. E. and Anderson, N. G., in: Radola, B. J. (Ed.), Electrophoresis 79, de Gruyter, Berlin 1980, pp. 329–339.
- [35] Taylor, J., Anderson, N. L. and Ancerson, N. G., in: Allen, R. C. and Arnaud, P. (Eds.), Electrophoresis '81, de Gruyter, Berlin 1981, pp. 383-400.
- [36] Anderson, N. L., Taylor, J., Scancora, A. E., Coulter, B. P. and Anderson, N. G., Clin. Chem. 1981, 27, 1807-1820.
- [37] Taylor, J., Anderson, N. L., Scandora, A. E., Jr., Willard, K. E. and Anderson, N. G., Clin. Chem. 1982, 28, 861-866.
- [38] Taylor, J., Anderson, N. L. and Anderson, N. G., Electrophoresis 1983, 4, 338-345.
- [39] Anderson, N. L. and Taylor, J., in: Proceedings of the Fourth Annual Conference and Exposition of the National Computer Graphics Association. Chicago, June 26–30, 1983, pp. 69–76.
- [40] Anderson, N. L., Hofmann, J.-P., Gemmell, A. and Taylor, J., Clin. Chem. 1984, 30, 2031-2036.
- [41] Anderson, L., in: Schafer-Nielsen, C. (Ed.), Electrophoresis '88, VCH Verlagsgesellschaft, Weinheim 1988, pp. 313-321.
- [42] Neidhardt, F. C., Appleby, D. A., Sankar, P., Hutton, M. E. and Phillips, T. A., Electrophoresis 1989, 10, 116-121.
- [43] Gil, G., Goldstein, J. L., Slaughter, C. A. and Brown, M. S., J. Biol. Chem. 1986, 261, 3710-3716.

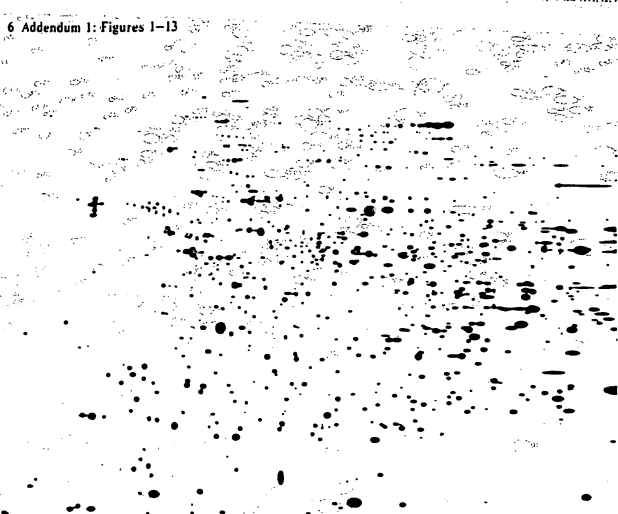


Figure 1. Synthetic representation of the standard rat liver 2-D master pattern, rendered as a greyscale image using a videoprinter.

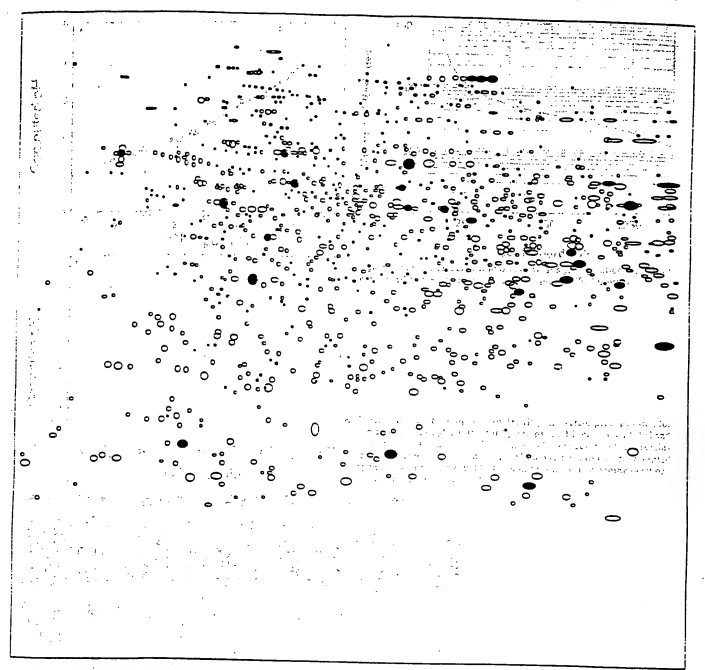


Figure 2. Schematic representation of the master pattern (the same as Fig. 1), useful as an aid in relating specific areas of Fig. 1 and the following detailed quadrants.

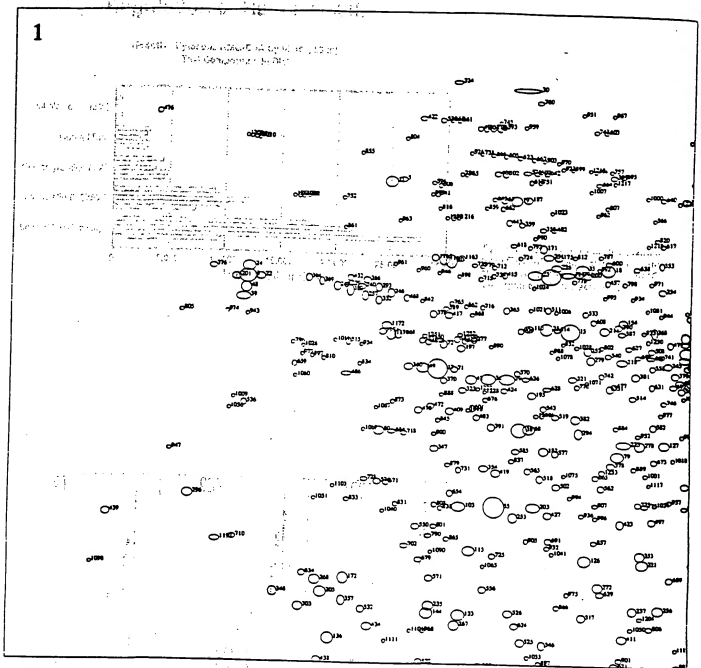


Figure 3. Upper left (high molecular weight, acidic) quadrant (#1) of the rat liver map, showing spot numbers.

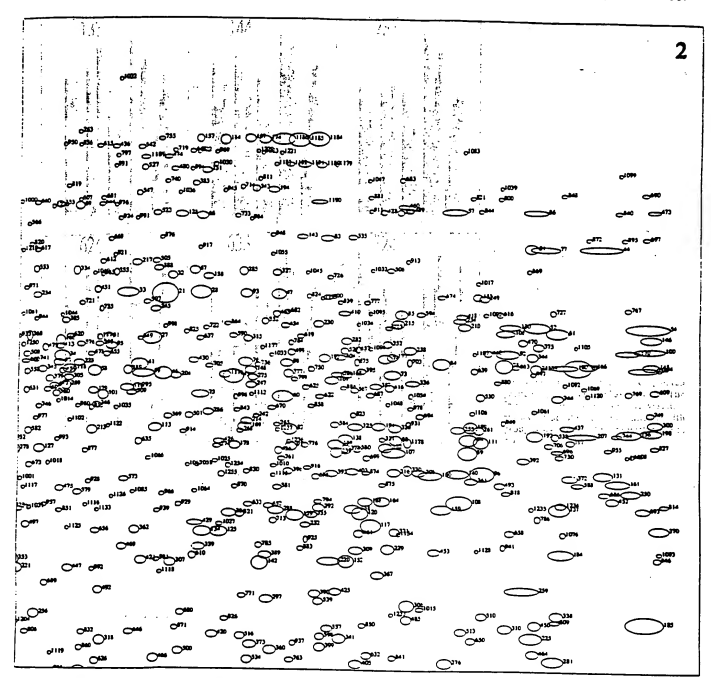


Figure 4. Upper right (high molecular weight, basic) quadrant (#2) of the rat liver map, showing spot numbers.

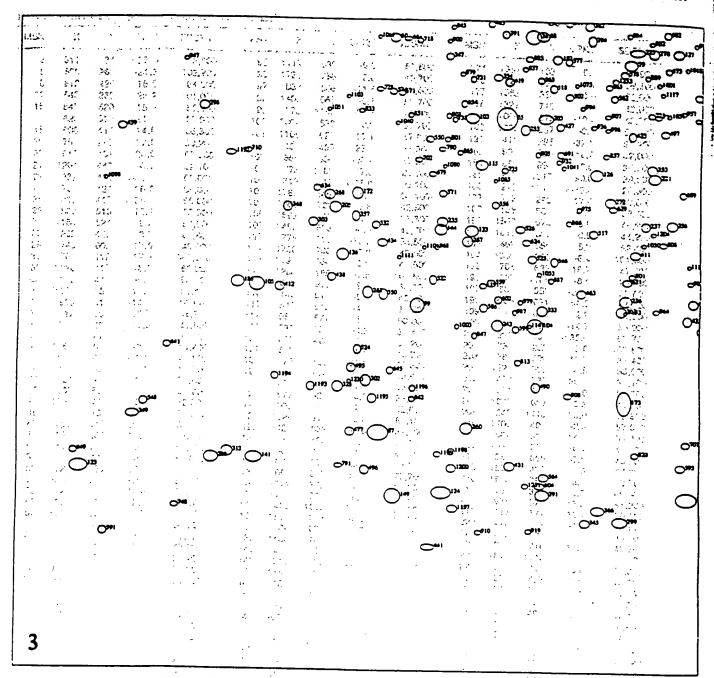


Figure 5. Lower left (low molecular weight, acidic) quadrant (#3) of the rat liver map, showing spot numbers.

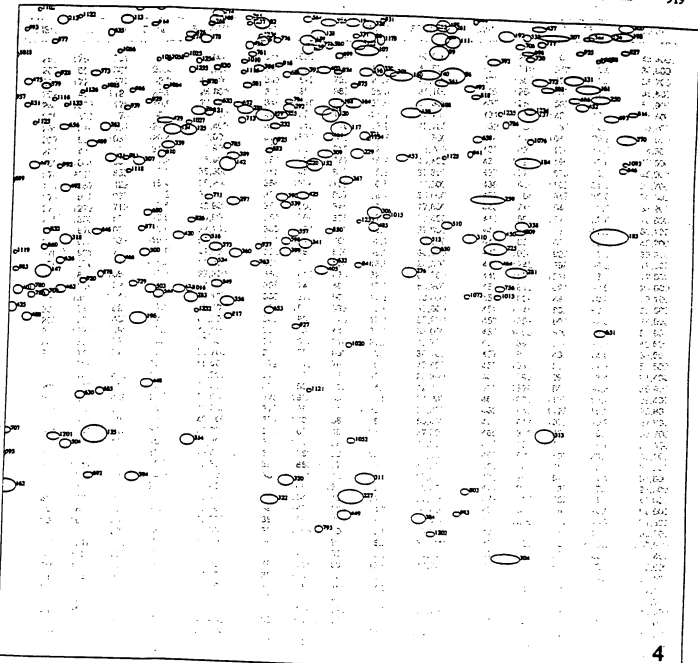


Figure 6. Lower right (low molecular weight, basic) quadrant (#4) of the rat liver map, showing spot numbers.

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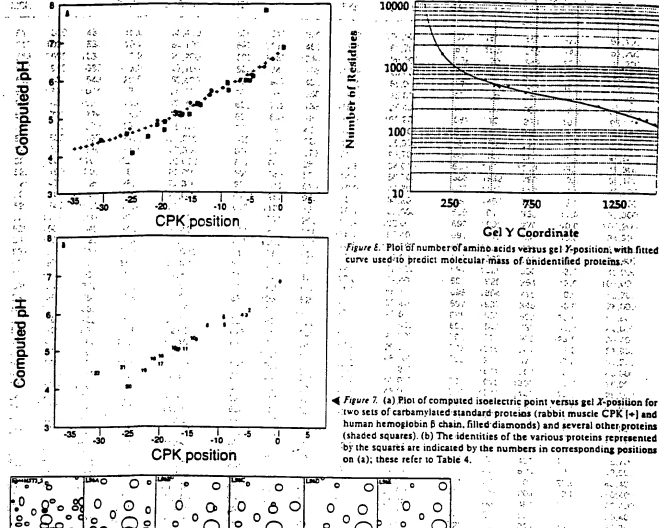
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Figure 9. Montage showing effects in the region of MSN:413. The montage shows a small window into one portion of the 2-D pattern, one row of windows for each experimental group, and one panel for each gel in the experiment. The left-most pattern in each row is a group-specific copy of the master pattern followed by the patterns for the five individual rats in the group. The highlighted protein spots (filled circles) are spot 413 (on the right of each panel, identified as cytosolic HMG-CoA synthase) and two modified forms of it (1250 and 933). From the top, the rows (experimental groups) are: high cholesterol, controls, cholestyramine, lovastatin, and lovastatin plus cholestyramine.

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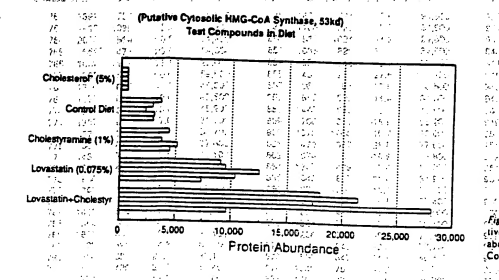
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Regulation of Rat Liver 413



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Figure 10. Bargraph showing the quantitaclive effects of various treatments on the abundance of MSN:413 (cytosolic HMG-CoA synthase) in the gels of Fig. 9.

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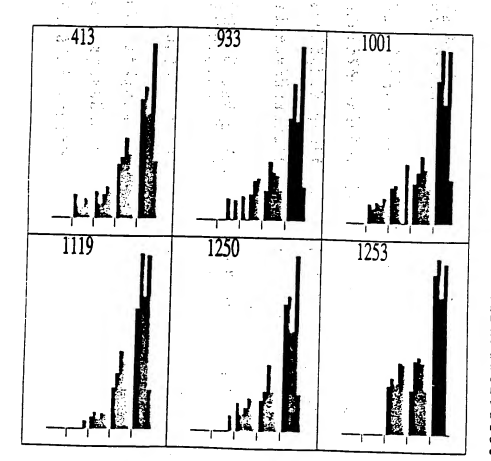


Figure 11. Bargraphs of a series of six coregulated spots including MSN:413. In the bargraphs, the abundances of the appropriate spot (master spot number shown at the top of the panel) in each animal are shown. The five five-animal groups are in the order (left to right); high cholesterol, controls, cholestyramine, lovastatin, and lovastatin plus cholestyramine. Each bar within a group represents one experimental animal liver (one 2-D gel). Note the correlated expression of the 6 spots, especially in the two far right (most strongly induced) groups.

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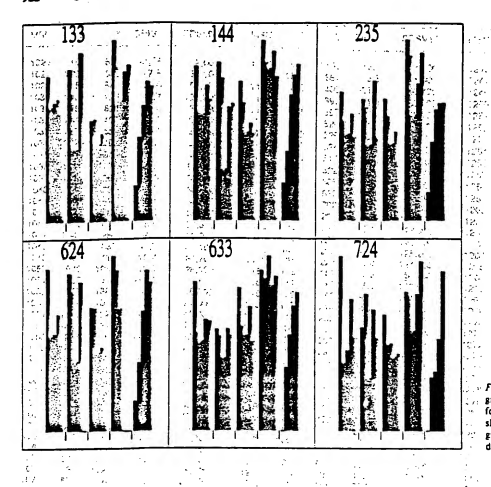


Figure 12. Data on a second coregulated group of spots, presented as in Fig. 11. The fourth experimental group (lovastatin) shows a modest induction, while the fifth group (lovastatin plus cholestyramine) does-not. 26 •

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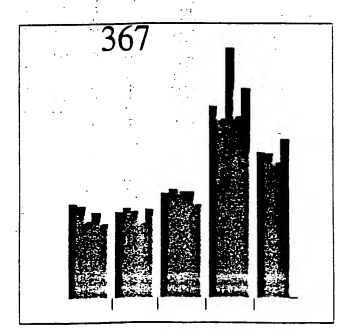


Figure 13. Data on spot MSN:367, presented as in Fig. 11. This protein shows unambiguously the anti-synergistic effect of lovastatin and cholestyramine (fifth group) as compared to lovastatin (fourth group). This response contrasts strongly with the regulation pattern seen in Fig. 11.

Table 1. Master table of proteins in the rat liver database"

ASN	,	į _C Y	CPKol	WikeCe	MS	N,	K	Y : CFKol	SDSMW	MSI	, N	Y	CPKel:	68615
3	311		ૄ~-35.0		<u> </u>	5	- 53	6 - 5.9	53,800	17				
5	568		-24.3	102,900	,*	6, . 173			40,700	17			-6.7	162,90
11	E12	4	-16.0 -25.2	64, 800 101,000	2	7 103			51,600	17			-15.7 -3.6	69,30
15	845		-15.3	55,200	•	8 140 9 57			51,700	. ; 17			-7.2	52,80 43,00
17	629		-21.6	50,000	. 10			,	25,000	17			-10.4	48,30
18	906		√ -14.0	66,300	10		.,		53,700 47,900	180		·, •••	-0.5	51,600
19	755		-17.5	90,200	່ 10		,		61,300	181 182		295	-32.1	91,200
20	649		20.9	67,900	10		630		37,300	18		730	-16.2	42,000
21 22	1204 332		* -8.7 <-35.0	62,100 63,800	10			17.0	23,800	185		1017	-0.6 [^] >0.0	34,500
23	787	424	-16.6	65,000	10: 10:		1117		2€,100	186	7	1113	<-35.0	29,600 26,300
24	313	417	<-35.0	66,000	10				56,100	187		296	-17.0	90,800
25	807	516	-16.1	55,500	108		720		42.500	-188		807	4.2	38,400
27	1184	524	-9.0	54,900	109		:593	4.8	38,300 49,700	. (191 192		674	•3.9÷	44,900
28 29	1263 743	446	-8.0	62,400	110		g '516		55.500	193		687 555	-0.9	44,200
30	768	605	-17.8 -17.2	49,000 348,600	111		700	2.0	43,500	7 194			-5.0 -6.4	52,400
32	1216	417	-8.6	€€,000	. 113 114		680	-8.9	44,500	195		632	-16.7	101,600 47,300
33	1145	445	-9.5	62,500	115		185 907	•7.5	160,800	196		1185	-8.4 ²	23,700
34	1037	555	-11.3	52,400	116		610,	-19.6 -9.5	34,100	197	_	553	-20.1	52,600
35	863	412	-14.9	66,600	117	1548	849	. ~0.5 ~4.1	48,700 / 36,500	∷"198. 199		C 681	>0.0	44,500
36 38	712 763	606	-18.7 -17.3	48,900 43,900	118	1050	577	-11.1	50,800	200	1711 872	674 424	-2.2	44,900
3 9	304	694 470	-17.3 <-35.0	43,800 59,800	120 121	1530	828	∢.3	37,400	201	292	435	<-35.0	65,000
41	1165	560	-9.2	51,400	121	638 1572	423	-15.4	65,200	202	736	253	-18.0	63,700 107,800
42	684	607	-19.6	48,800	123	23	712 1433	-3.8 <-35.0	42,900	203	786	829	-16.7	37,400
43	1318	589	-7.3	50,000	124	621	1474	·21.9	15,300 <u>1</u> 13,900	204	1224	589	-8.5	50,000
44 45	1924 1203	362 586	-0.1	74,600	125	1298	862	-7.5		205	439 1994	983	-30.9	31,100
47	1391	447	-8.7 -6.3	50,200 €2,300	126	672	921	-14.7	35,000 · · · · · · · · · · · · · · · · · ·	207	1895	571 687	- >0.0 0.3	ु:51,300
18	309	454	<-35.0	€1,500	127 128	1000 1229	717	-12.0	42,60C	208	240	1418	<-35.0	15,800
19	605	587	-22.5	50,100	129	1422	311 832	-8.4	86,10C	. 210	1700		-23	57,000
0	621	535	-21.8	53,900	130	1776	499	-5.8 -1.4	37,300 57,000	211	902	517	-14.1	55,400
1 2	1113	522	-10.0	55,000	131	1930	757	-0.1	40,700	213 1214	1087 1340	684	-10.4	44,400
3	1620 725	499 177	-0.9 -18.3	57,000 170,800	132	660	537	-20.4	53,800	215	1591	668 495	-7.0	: 45,200
4	2001	500	>0.0	56,900	133 134	666	1019	-20.2	29,700	216	1585	755	-3.5 -3.6	57,300 40,700
5	722	830	-18.4	37,300	135	1271 1161	862 1389	-7.9	36,000	217	1159	393	9.3	69,300
56	€78	. 533	-19.8	54,100	136	453	1063	-9.3 -29.7	16,800 28,100	218	931	572	-13.5	51,200
7 8	1682	302	-2.5	89,000	137	1858	823	-0.6	37,70C	219	713 14 79	177	-18.7	170,500
9	1091 1171	580 585	-10.3 -9.2	50,600	138	1504	697	4.6	43,700	221	965	911 927	-4.9 -12.8	33,900
io .	1400	624	-6.2	50,300 47,800	139 140	1488 1689	707	4.8	43,200	223	934	716	·12.6	33,300 42,700
1	1853	508	-0.6	56,200	141	311	756 1417	-2.4	40,700	225	1812	1045	-1.0	28,800
2	1888	567	-0.4	51,500	142	1366	915	<-35.0 -6.7	15,800 33,800	226	821	411	-15.8	66,800
5	735	297	-18.1	90,500	143	1429	346	-5.7	77.900	227 228	1586 1065	1483	-3.6	13,600
	1263 1252	312 407	-8.0	85, 900	. 144	615	1017	-22.1	29,800	229	1577	567 890	-10.8	51,600
B	779	692	-8.1 -16.8	67,300 43,900	145	2006	566	>0.0	51,600	230	1458	496	-3.7 -5.2	34,800
	1064	296	-10.8	90,800	146 147	2006 1070	518 1108	>0.0	55,300	232	1440	849	-5.5	57,300 36,500
1	656	589	-20.6	50,000	148	1347	578	-10.7 . -6.9	26,500	234	1692	489	-2.4	57,900
?	638	545	-21.2	53,100	149	541	1481	-25.7	50,800 13,700	235 236	618	1004	-22.0	30,300
	1582	583	-3.6	50,400	150	1645	760	-2.8	40,500	237	920 952	1138	-13.7	25,400
	1570 1264	556 621	-3.8 -8.0	52,300 48,000	151	1269	236	-7.9	117,000	238	1611	1008 541	∙13.1 •3.2 _. .	30,200
	1338	564	-7.0	51,800	152 153	1507	911	4.5	33,900	239	1489	720	-4.8	53,500 42,500
	1833	363	-0.8	74,400	154	1722 932	448 503	-2.1	62,100	240	501	448	-27.7	62,100
	1767	565	-1.5	51,700	155	1031	294	-13.5 -11.4	56,600 91,400	241	1820	569	-0.9	51,400
)	925		-13.6	41,600	156	1970	684	>0.0	44,400	242 243	1357 711	658	-6.8	45,800
١.	534 1811	698	-26.1	43,600	157	1258	183	-8.1	162,400	244	1855	1182 621	-18.7	23,800
	412	363 681	-1.0 -6.0	74,500 44,500	· 158	1275	417	-7.8	65,900	245	1189	474	-0.6 -8.9	48,000
	471	347	-5.0	77,500	159 160	1663 1034	820	-2.6	37,800	246	551	459	·25.1	59, 300 61,000
	662	563	-2.7	51,800	161	1953	527 771	-11.4	54,600	247	1348	604	-6.9	49,100
	596	479	-3.4	5E,900	162		1482	>0.0 -11.6	40,000 13,700	248	460	448	·29.3	62,100
	817	301	-0.9	89,100	164	1566	606	-3.8	38,400	249 250	1733 1974	451 788	-1.9	61,800
			-27.0	17,400		1905	565	-0.2	51,700	250 251	808	788 392	>0.0	39,200
	589 706	698 719	-3.5 -2.2	43,600 42,500		1340	181	-7.0	164,900	252	874	553	-16.1 -14.6	69,500 53,500
	651		20.8	42,500 81,700		1506 1338	583 578	-4.6	50,400	253	753	848	-17.6	52,500 36,500
	415	710	-6.0	43,000		1969	678 541	-7.0 >0.0	44,700	254	995	450	-12.1	61,900
1	773	545	-1.4	53,200	171	800	378	>0.0 -16.3	53,500 21,800	255	1690	679	2.4	44,600
	338 708	446	-7.0	62,300	172	476	958	·28.7	71,800 32,100	256 257	994 508	1006	-12.1	30,200
		696	-2.2	43,700	173	919	314	-13.7	,	£3/	340	464	-27.4	60,400

Master table of proteins in the rat liver database, showing spot master number, gel position (x and y), isoelectric point relative to CPK standards, and predicted molecular mass (from the standard curve of Fig. 8).

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MSN	×	Y	CPKol	SDSMW	MSN	X	Y	CPKol	SDSMW	MSN	×	Y	CPKol	SDSMW
259	1796	e49617	-1.1	31,900		1006	578	-11.9	50,800	426	1296	704	-7.6	43,300
260	661	1361	-20.4	17,700	346	1095	640	-10.3	46,800	-427	810	843	-16.0	36,800
261	1725		-20	24,600	2:347	; ÷ 625	728	-21.7	42,000	428	1565	303	3.9	86,700
262	496	1127	-28.0	25,800 177,400	2 348 349		1343	35.3	31,100 · · · · · · · · · · · · · · · · · ·	429 430	1259	847	-8.0	36,600
:263	1063	172 673	-10. 9 -6.3	45,000	350	² 521	1130	`<-35.0 26.7	25,700 5	431	1253 734	562 1426	-8.1 -18.1	51,900
(265 √266	1390 510	437	-27.3	63,400	5° 351	912	€19	-13.9	48,100	432	483	433	-28.5	15,500 63,900
267	560	1038	-20.4	29,000	₹ 352	2/1574	530	3.7	54,300	434	518	1041	-26.9	28,900
268	430	961	-31.0	31,900	ं 353		S12	-12.9	33,900	435	1020	1170	-11.6	24,300
269	1044	606	-11.2	4E,900	≳(354		7€2	-18.9	40,400	436	1122	196	-9.8	147,600
270	2019	853	>0.0	3€,300]r 355	g:1450	630	-5.3	37,300	437	1870	673	-0.5	45,000
271	857	422	-15.0 -14.2	65,200 31,700	∂ 356	~1374	1152	-€.5	24,900 ° 30,600 °	438	435	1102	-31.0	2€,700
272	895	968 712	-7.6	42,900	:: 357 :: 358	798	997 346	-28.7 -16.3	30.600 S	439 440	86 1740	847 544	<·35.0	3€,600
275	1292 1350	590	-6.9	49,900	359	764	338	-17.3	79.400 7h	441	599	1571	-1.8 -22.8	53, 200 10, 800
1 276	1670	1089	-2.6	27,100	360	1354	1068	₹ -6.4	27,900	443	743	335	-17.8	80,100
277	688	538	-19.4	53,700	361	1713	769	-2.1	40,100 50	446	801	668	-16.2	45,200
278	961	718	-13.0	42,600		≟-1161	85 9	-9.3	36,100 ∂ €	447	1050	926	-11.1	33,300
279	£79	570	-14.5	51,300	44	3/ 914:	1156	-13.B	24,800	448	1245	1298	-8.2	19,800
281	1648	1084 525	-0.7 -4.6	27,300 54,800	% 364 5 365	? 412° "> 741.	435; 486;	32.0	63,700	449	1576	1516	-3.7	12,600
282 283	1505 1313	1147	-7.3	25,100	366	£78	1503	-17.9 -14.6	58,200 13,000	450 451	1818 1094	1021 440	-0.9	29,600
284	1314	829	-7.3	37,400	367	1560	935	-3.9	33.000	452	1945	802	-10.3 >0.0	63,100 38,600
: 285	1332	408	-7.1	67,200	₂ , 368	983	520	-12.4	55,200	453	1652	894	-2.8	34,600
gn 286	1277	652	-7.8	46,100	369	434	441	-31.0	63,000	454	1403	500	-6.1	56,900
288	1391	824	-6.3	37,600	370 371	639	610	-21.2	48,700	456	1394	718	-6.3	42,600
289	1147 925	579 511	- 9 .5 -13.6	50,700 55,900	2 371 2 372	1587 1875	860 762	-3.6	36,100	457	905	436	-14.0	63,500
290 291	787	1476	-16.6	13,900	2 373.	. 1351	1059	-0.5 -6.8	40,400 28,300	450 460	1038 1598	581 294	-11.3 -3.4	50,500
262	1462	818	-5.1	37,800	374	1506	715	4.6	42,700	461	1528	863	-3.4 -4.3	91,400 35,900
293	531	449	-26.3	€2,000	375	1823	532	-0.9	54,200	462	1098	1137	-10.2	25,400
294	860	698	-14.9	43,600	376	254	417	<-35.0	65,900	463	849	1125	-15.2	25,800
295	1162	609	-9.3 25.0	48,700	377		583	-6.1	50,400	464	1814	1072	-0.9	27,800
296	218	814 979	<-35.0 -6.5	38,000 31, 30 0	378	621 1017	494	-21.8	57,500	465	1388	481	-6.3	58,700
297 299	1377 91 3	1523	-13.9	12,400	3/9	953	595 598	-11.7 -13.1	49,600 49,400	466 468	1194 577	1084 467	-8.9 22.0	27,300
300	2012	667	>0.0	45,300	382	856	674	-15.0	44,900	469	1140	888	·23.9 ·9.6	60,1 00 34, 900
301	702	178	-19.0	169,200	383	1252	258	-8.1	105,300	470	1797	524	-1.1	54,800
302	494	1280	-28.1	20,400	384	1699	1518	-2.3	12,500	471	1293	1133	-7.6	25,500
303	403	1008	-32.6 -0.7	30,100 10,300	385	1042	493	-1.1.2	57,500	472	618	655	-21.9	46,000
304	1843 1049	1585 593	-11.1	49,800		, 1490, 1554	583 603	-4.7 - 4.0	50,400 49,100	473 474	2009	299	>0.0	89,900
306	1608	989	-3.3	30,900	357 388	1193	404	-8.9	67,700	475	1205 1035	215 788	-8.7 -11.4	131,300 39,200
307	1219	916	-8.5	33,700	389	1374	902	-6.5	34,300	476	160	155	<∙35.0	207,600
306	1627	755	-3.0	40,700	390	1456	969	∙5.2	31,700	477	469	1370	-28.9	17,400
309	1524	892	4.4	34,700	391	718	690	-18.5	44,000	478	599	662	-22.8	45,600
310	1769	1028	-1.5 -3.3	29,400 14,700	392 393	1799 1482	732	-1.1	41,900	479	1009	540	-11.8	53,500
311 312	1609 266	1451 1408	<-35.0	16,100	394	1227	758 1461	-4.8 -8.4	40,600° 14,400	. 480 . 482	1216 816	235	-8.6	117,400
. 313	1902	1365	-0.3	17,600	395	1530	577	-4.3	50,800	483	693	346 673	-15.9 -19.3	77,800 44,900
214	1316	1395	-7.3	16,600	396	1410	755	-6.0	40,800	485	1608	1013	-3.3	30,000
315	1341	523	-7.0	54,900	397	912	256	-13.9	106,400	486	478	599	-28.6	49,300
318	1104	1053	-10.1	26,500	399	1465	1063	·5.0	28,100	. 487	1025	607	-11.5	48,800
320	1480 850	145 0 603	-4.9 -15.1	14,400 49,100	400 401	1473 1029	450	-4.9	61,900	488	1045	1186	-11,2	23,700
321 322	1454	1494	·5.3	13,300	403	1516	1140 754	-11.5 -4.4	25,300 40,800	489 - 490	1609 775	301 1289	-3.3	89,200
323	670	626	-20.0	47,700	404	1495	554	4.7	52,500	491	692	178	-17.0 -19.3	20,100 169,300
324	655	101	-20.6	420,500	405	1525	1092	-4.3	27,100	492	1100	964	-10.2	31,800
325	1521	675	4.4	44,800	406	723	252	-18.4	108,000	493	1760	776	-1.6	39,700
326	1587	677	-3.6	44,700	409	650	663	-20.8	45,500	494	882	247	-14.5	110,700
327	1388	409	-6.3 20.0	67,000	410	1501	478	-4.6	56,000	495	470	1258	-28.9	21,200
32 8 330	448 1608	1 <i>2</i> 91 751	-30.0 -3.3	20,100 40,900	411 412	936 350	1057 1120	-13.4 -35.0	28,300	496	494	1436	-28.1	15,200
330	1566	697	-3.8	43,700	413	1033	538	-35. 9 -11.4	26,000 53,700	497 499	980 1414	852 546	-12.5 -6.0	36,400 53,100
332	531	471	-26.3	59,600	415	737	425	18.0	64,900	500	1234	1072	-8.3	53,100 27,800
333	784	1156	-16.7	24,700	416	1578	606	-3.7	48,900	501	1246	659	-8.2	45,700
334	1059	407	-10.9	67,300	417	646	496	-21.0	57,300	502	824	792	-15.7	39,000
335	1593	303	-3.5	88,500	418	1695	482	.2.3	58,600	503	1246	1134	-8.2	25,500
336	1616	598	-3.2 -0. 6	49,400 30,300	419	725	770	-18.3	40,000	504	1115	1407	-9.9	16,200
338 339	1854 1265	1004 888	-0.6 -8.0	30,3 00 34, 900	420 421	1289 1171	1041 912	-7.7 -0.1	28,900	505 506	1189	391	-8.9	69,700
340	581	585	-23.6	50,300	422	599	162	-9.1 -22.8	33,900 193,700	507	1578 787	402 250	-3.7 -16.6	68,000
341	1497	1047	4.7	26,700	423	929	856	-13.6	36,200	508	979	552	-10.6	109, 000 52, 600
343	1351	265	-6.8	102,200	424	739	625	-17.9	47,700	509	1153	619	-9.4	48,100
344	1813	549	-0.9	52.800	425	1490	965	-4.7	31.800	510	1730	1006	-2.0	30.200

	· · · ·	137, 19	CPKel	SC. SDSMW	-: : MS/ - —	t e d	Υ.	CPKol	SDSMW	• .	WZN	, X	<u> </u>	CPKel	SDSWW
511 512	806 1096			56,400 54,100	590 561			-21.9	100,500	riki. B	674	1661	448	-27	; 62,100
513	1696		2 200	25,200	590			-9.1 -5.0	60,700 28,800			1523	. 562	- 44	51,900
514	54	,		47,100	596		1188	-17.9	23,600	2	: 676, : 677	708 919	642,	-18:8	46,700
515	481			53,400	600		402	-14.0	68,000		678	1085	615 551	-13.7 -10.5	, 48,300 52,700
516	1334 868			28,800	601		658	-19.5	45,800		679	600	4 923	-22.7	33,400
517 · 518	798			29,700 39,600	602		1138	-18.7	25,400		680		€ 1004°	-8.3	30,300
519	822			45,100	604			-14.1	165,200	7			· 283		95,100
520		-		189,000	605		1461 223	-16.7 -18.0	14,400	3	682		477		59,100
521	. 1332			37,300	606	£29	273	-21.6	125,300 98,700		683 684	1596. 555	249	-3.4.	109,800
522	603			2€,600	607	1064			್:⊶.‱	42		1167	699 1 1313	-24.8 -9.2	43,500
523 ; 524	1190 479	309 1226		86,800	608	883	503	-14.5	5E,700	3		1932		0.0	15,300 39,100
525	768	1066	-17.2	22,300 25,000	609 610	2012 1255	610	>0.0	48,700	•	687 .	1545	61913		548,100
526	747	1016		25,800	612	1103	903 391	-8.1	34,200	; ;		1456	764	-5.2	40;300
527	1170	231	-9.2	119,600	£13	778	265	-10.1 . -16.9	69,600		689	1011	953	-11.8	32,300
28	1502	542	4.6	53,400	614	·E24	518	-15.7	102,000 55,400	,	690 691	1995 812	270	>0:0	100,200
30	1728	- 620	∺ -2.0	48,000	615	1095	195		149,100	ż			888'' 1461'	-16.0 -9.4	34,900 14,400
32	507	1011	-27.4	30,000	616	1759	478	-1.6	59,000	۷,		1993	819:5	>0.0	14(400) 37,800 1
33 34	870 1347	489 1085	-14.7 -6.9	57,900 27,300	617 618	994 751	372	-12.1	72,900		694	1628	656	-3.0	45,900
35	1513	346	4.5	77,800	619	751 1429	374 518	-17.6	, 72,400	.~1	695	928	254)	-13.6	107,000
36	306	654	<-35.0	46,000	€20	1050	520	-5.7 -11.1	_55,300	5.	696	1854	715	-0.6	42,700
38	1851	689	-0.7	44,100	€21	923	1105	-13.7	55,200 26,600	۲.,	697 698	1997 957	345	>0.0	78,000
39 40	1463	982	· -5.1	31,100	€22	1462	€22	-5.1	47,900	<i>)</i> .			563 € 730 €	-13.0 -4.2	51,800 42,000
11	90 9 625	561 289	-13.9 -21.7	52,000 93,100	€23 €24	759	225	-17.4	124,000	4	702	577	900	-23.8	34,400
12	1164	198	-9.2	146,200	625	758 1438	1038 606	-17.4	29,000	٠.	703	1610	562	-3.2.	51,900
ß,	803	655	-16.2	45,900	€26	1096	1069	-5.5 -10.2	48,900		705	1278	571	-7.8	51,200
4	1259	1143	-8.0	25,200	€27	542	548	-13.3	27, 20 0 53,000		706 707	1841	704	-0.7	43,300
15	856	1526	-15.0	12,200	€28	809	,621	-16.0	48,000	٠, ٠	709	1018	.1386 ∂1145: /	-11.7 -10:7	16,900
15 17	803 1162	1071 274	-16.2 -9.3	27,800 96,400	£29	699	979	-14.1	31,300	•	710	293	. 889	<-35.0	25,100 . 34,800
8	128	1321	<-35.0	19,000	630 631	1135 97 9	1321	-9.6	19,100		712	720	412	-18.5	66,600
9	1355	1122	-6.8	25,900	632	1542	615 1076	-12.5 -4.1	48,300		713	1386	841	-6.4	36,800
io	595	866	-23.0	35,800	633	1345	814	-6.9	27, 600 38, 00 0		714 715	1328	263	-7.1	103,100
2	1369	494	-6.6	57,500	634	409	950	-32.2	32,400		716	698 701	433 481	-19.1	63,900
3 5	992 1125	405 410	-12.2 -9.8	67, 600	635	1165	704	-9.2	43,300		717	1875	699	-19.0 -0.5	58,700 43,600
6	705	975	-18.9	66, 900 31, 400	636 637	774 1263	604	-17.0	49,000		718	575	702	-23.9	43,400
7	1477	1030	4.9	29,300	638	952	524 411	-8.0 -13.1	54,800 66.700		719	1216	204	-8.6	140,400
8	980	583	-12.5	50,400	639	1717	575	-2.1	66,700 51,000		721 7 <u>22</u>	1069 1272	454	-10.8	60,400
9	700	1109	-19.1	26,400	640	994	292	-12.1	92,000		723	958	506 822	-7.9 -13.0	56,400
0 2	1028 898	621 794	-11.5	4E,000	641	165	1224	<-35.0	22,400		724	763	395	-17.3	37,700 69,100
1	789	1446	-14,1 -16.6	38,900 14,900	642 643	803	251	-16.2	106,900		725	720	916	-18.5	33,700
5	777	766	-16.9	40,200	644	719 1100	296 294	-18.5	90,700		726	1476	415	4.9	66,200
5	980	328	-12.5	81,900	645	534	1263	-10.2 -26.1	91,400 21,000		727 728	1846	473	-0.7	59,400
	1519	611	-4.4	48,600	646	1153	1038	-9.4	29,000		728 729	510 1217	783 1126	-27.3	39,400
	1212	661	-8.6	45,600	648	1246	204	-8.2	140,000		730	1858	724	-8.6 -0.6	25,800 42,300
) 	760 618	594 956	-17.4 -21. 9	49,700 32,100	649 660	14		<-35.0	16,200		731	665	765	-20.2	42,300 40,300
	1142	771	-9.6	40,000	650 651	1713 1986	1049 1183	-2.1 -0.0	28,600		733	1321	312	-7.2	85,900
ı	532	787	-26.2	39,300	652	1378	816	>0.0 -6.5	23,800 38,000		734	719	427	-18.5	64,600
;	771	250	-17.1	109,200	653	1442	1165	-c.5 -5.5	24,400		_	1101 1359	473 560	-10.2	59,500
	1068	534	-10.8	54,100	654	650	806	-20.8	38,400		738	696	569 220	-6.7 -19.2	51,400 127,600
	822	734 754	-15.7	41,800	655	1111	551	-10.0	52,700		739	687	409	-19.2	127, 600 67, 000
	914 1064	754 794	-13.8 -10.8	40,800 38,900	656 657	1095	861	-10.3	36,000			1205	256	-8.7	106,200
	1524	714	-10.8 -4.4	42,800	657 6 5 8	1524 1777	540 860	-4.4	53,600		741	995	563	-12.1	51,900
	392	783	-6.3	39,400	659	391	584	-1.4 -33.4	36,000 50,400		742	998	596	-14.1	49,500
	962	686	-12.4	44,200	660	977	565	-12.5	50,400 51,700		743 744	881 1951	181 686	-14.5	165,900
	487	672	4.8	45,000	661	€58	166	-20.5	187,500		745	726	686 168	>0.0 -18.3	44,200 183,600
	758 687	731 1152	-17.4 -10.5	41,900	662	732	312	-18.1	86,100		746	999	643	-12.0	46,600
	930	523	-19.5 -13.5	24,900 55,000	663 664	1787	567 250	-1.2	51,500		748	182		<-35.0	13,000
	888	774	-13.5 -0.4	39,900	664 665	888 889	268 775	-14.4	100,900		_	2005	649	>0.0	46,300
	642	485	-21.1	58,300	666	715	775 221	-14.3 -18.6	39,800			1448	575	-5.4	51,000
	317	519	-7.3	55,300	667	781	227	-16.8	126,300 122,400		751 752	792 450	266	-16.5	101,900
			c-35.0	11,500	668	646	165	-21.0	189,100		752 754	469 664	296 254	-28.9	90,600
	014	614	-11.7	48,400	669	1116	353	-9.9	76,300		_	1195	184	-20.3 -8.8	107,000
	732 527	176 478		172,300		1382	643	-6.4	46,600				1113	-0.9	161,000 26,300
- 10	627	478	-3.0	59,000	671	547	789	-25.3	39,200		757	909	246	-13.9	111,000
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· ' ;	MSN	X	. Y	CFKol	SDSMW	MSA	х,	(Υ.	CFKol ;	SDSWW	MŚN	X	\$ 'Y	"CPKol"	SOSWW
ا ۇ ئ	761	1390	733	6.2	41,800	64		271	-0.6	99,500	£39	1197.	. 827	-8.8	37,500
ş 7.	763 . 764	1416	1085 569	5.9 .>0.0	27,300 51,400	54) E54		523 _. 1024	-5.2 -4.2	54,900 29,600	941 642	1765	885 472	-1.5 -22.7	35,000 50,600
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	768 769	1330 1570	685 613	-7.1 >0.0	44,300 48,500	. E50		194 890	-10.9 -14.4	150,500. 34,800	946 947	630 187	423 736	-21.6 <-35.0	65,100 41,600
	,770	857	617 ·	-15.0 -7.0	4E,200	858 859		639	-5.4	4E,900	948	1380	> 344	-6.5	7E.200
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	775 776	969 1438	706	-12.8 5.5	37,600 43,100	861 862	472 674.	347 480	-28.8 -19.9	77,500 58,800	951 952	960 957	152 701	-14.9 -13.0	213,000 43,400
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	778 779	850 700	~-411 :	-19.1	66,800	866	827.	z 1004	-21.0 15.€	34,900 30,300	955° 957	1938 1010	712 816	>0.0 -11.8	42,900 37,900
	780 784	1052 1413	1136 529	-11.1 -6.0	25,500 54,400	668 669	685 1807	494 402	19.5 -1.0	57,400 68,000	959 960	768 596	174 419	-17.2 23.0	174,900 65,700
	785 786	1364 1822	885 835	-6.7 -0.9	35,000 37,100	670 871	1323	783 1031	-7.2 -€.4	39,400 29,300	961 962	557 887	409	24.8	67,100
	767	563	392	-14.3	69,500	٤72	1904	346	-C.3	77,700	963	564	320 334	-14.4 -24.5	83,900 ···. 80,500 ···.
;	790 791	616 451	882 - 1429	-22.0 -29.8	35,100 (15,400	E73 6 5 ° €74	556 11540	647 756	-24.5 -4.2	46,400 40,700	964 965	969 671	1155 255	-12.8 -20.0	24,800 106,600
1,57	792 763	777 1536	377 1543	-16.9 -4.2	72,000 11,700	675 676	1566 1198	777 351	-3.8 -8.8	39,700	966	1204	796	-8 .7	38,700
	. 794	1461	807	-5.1	38,300	έ77	1076	720	-10.6	76,800 42,500	967 968	910 609	154 1048	-13.9 -22.3	210,300 28,700
	796 757	388 11 2 6	546 212	-33.6 -9.8	53,100 133,700	E78 E79	1161 647	1111 757	-9.3 -20.9	26,400 40,700	969 970	1285 822	206 232	-7.7 -15.8	138,900 7 / 119,300
- 1	798 799	933 1420	437 593	-13.5 -5.9	63,400 49,800	850 861	1756 1543	594 278	-1.6 -4.1	49,700.	971 972	976	437.	-12.6	63,400
	600	1750	279	-1.6	9€,500	663	1432	890	-5.7	97,100 34,800	974	403 279	567 495	-32.6 <-35.0	51,600 · · · · 57,400 · · · ·
	801 802	624 898	865 547	-21.7 -14.2	35, 800 53, 000	654 655	922 1103	689 414	-13.7 -10.1	44,100 66,400	975 976	844 1124	981 295	-15.3 -9.8	31,200 91,100
	803	1775 573	1468 196	-1.4 -24.0	14,200 1 148,400 7	856 887	1501 798	607 1103	-4.6 -16.3	48,900 26,600	977 97 8	994 1612	664	-12.1 -3.2	45,400
	805 506	203 980	494 1039	<-35.0 -12.5	57,400 29,000	888 689	636	634	21.3	47,200	979	749	1141	-17.7	46,700 25,300
	807	902	308	-14.1	E7,200 ·	890	9 51 717	759 548	-13.1 -18.6	40,600 52,900	980 981	1064 1197	642 911	-10.8 -8.8	46,700 ≥ 33,900
	808 809	625 1851	827 1015	-21.7 -0.7	37, 500 29,900	891 892	1123 891	229 413	9.8 -14.3	121,200 65,400	983 984	1762 1344	1508 317	-1.6 -6.9	12,800 84,700
	810 811	440 1358	573 249	-30.9 -6.8	51,100 109,700	894 895	1245 1962	234 346	-8.2 >0.0	117,800 77,700	985 987	1024 739	1105 1159	-11.5	26,600
	£12	851	393	-15.1	69,400	596	1322	626	-7.2	47,700	988	816	555	-17.9 -15.9	24,600 52,400
	813 814	745 2028	1246 810	-17.8 >0.0	21,600 38,200	897 898	420 662	570 428	-31.4 -20.3	51,300 64,500	990 3 6 1 991	785 1159	361 317	-16.7 -9.3	74,900 84,500
	815 816	1086 629	645 313	-10.4 -21.6	46,500 85,700	899 900	845 624	243 703	-15.3 -21.7	113,000 43,400	992 993	1090 1030	928 701	-10.4 -11.5	33,300 43,400
	817 618	1376 1771	1177 790	-6.5 -1.4	24,000 39,100	901 903	931 799	1094	-13.5	27,000	994	847	811	15.2 ₁	38,200
	819	1045	263	-11.2	103,100	904	765	229 520	-16.3 -17.2	121,000 55, 20 0	995 996	902 888	461 847	14.1 -14.4	60,7 00 36,6 00
	E20 621	984 1712	362 279	-12.4 -2.2	74,600 96,700	905 907	775 888	889 824	-17.0 -14.4	34,800 37,600	997 998	1815 ¹ 1205	579 504	-0.9 -8.7	50,700 56,500
	822 823	1256 1517	205 654	-8.1 -4.4	139,200 46,000	906 910	828 681	1303 1544	-15.6 -19.7	19,700 11,700	999 1000	- 617	289	-22.0	93,100
	E24	1442	449	-5.5	62,000	911	1544	301	-4.1	85,100	1001	968 970	290 771	-12.8 -12.7	92,700 40, 000
	825 826	1240 1309	513 1014	-8.3 -7.4	55,800 29,900	913 914	1606 1237	387 688	·3.3 -8.3	70,400 44,100	1002 1003	1736 . 643	478 1184	-1.9 -21.1	58, 900 23,7 00
	827 828	2012 937	708 1405	>0.0 -13.4	43,100 16,200	916 917	1442 1260	~ 749 367	-5.5 -8.0	41,100 73,700	1006 1007	822 875	487	-15.8	58,100
	830	1342	756	-7.0	40,700	919	764	1541	-17.3	11,700	1009	291	279 644	-14.6 <-35.0	96,400 46,600
	831 832	562 1073	826 1 039	-24.5 -10.7	37,500 29,000	920 921	1133 1123	1123 380	9.7 -9.8	25,900 71,500	1010 1011	1386 459	745 541	-6.4 -29.4	41,200 53,500
	833 834	481 501	820 581	-28.5 -27.8	37,800 50,500	923 924	629 1131	242 318	-15.6 -9.7	113,200 84,300	1012 1013	€79	661	-19.7	45,600
	837	751	748	-17.6	41,100	925	1441	- 874	-5.5	35,400	1014	1818 1032	1128 634	-0.9 -11.4	25,800 47, 200
	838 839	635 14 94	833 459	-21.3 -4.7	37, 200 60,900	926 927	679 1487	219 1191	-19.7 -4.8	128,200 23,500	1015 1016	1629 1311	994 1134	-3.0 -7.4	30,700 25, 500
	840 841	1952 1585	301 1080	>0.0 -3.6	89, 300 27, 500	928 929	1082 1231	775 816	-10.5	39,800	1017	1722	424	-2.0	65,000
	842	571	1312	-24.1	19,400	931	1609	670	-8.4 -3.3	38,000 45,100	1018 1020	1015 1574	743 1219	-11.7 -3.7	41,300 22,500
	643 644	1325 1727	649 301	-7.2 -2.0	46,300 89,200	632 933	810 965	900 520	-16.0 -12.8	34,400 55,100	1021	781 1129	484 83	-16.8 -9.7	58,400 591,300
	845	630 2016	679 905	-21.5 >0.0	44,600 34,200	934 936	947 865	462 843	-13.2	60,600	1023	812	317	-15.9	84,600
	847	673	1200	-19.9	23,200	937	1421	1056	-14.8 -5.9	35,800 28,400	1024 1025	785 1290	446 739	-16.7 -7.7	62,400 41,500

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1026	405	532	-32.5	52,600	115			9 -13,7	24,700	1246	547	577 .	-25.3	***
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1030	. 1264	.· 226	7.7	123,200	116				6E,400	1249	516	572	-27.0	51,200
I C31	\$86	622	-12.3	37,700	116	3 66			- 6E,800 6E,700	1250 1251	973 607	536 532	-12.7	53,900
1633	1547 1381	403 551	-4.1 -6.4	€7,900 52,700	116			-24.4	54,500	1252	665	529	-22.4 -20.2	54,200 54,400
634	1525	496	4.3	57,200	117 117				54,500	1253	899	766	-14.1	40,200
C35	1128	645	-9.7	4E,500	117				54,800 55,700	1254 1255	1311	746	-7.4	41,200
035	1226	274	-8.5	9E,300	- 1174			-10.2	55,000	1255	1300 1938	761 712	-7.5 0.0	40,400
649 640	1761 541	262 839	-1.6 -25.7	103,600 36,900	1170				5C,200	1258	1806	718	-1.0	42,900 42,600
041	€18	S10	-15.8	34,000	1178				53,700	1259	1727	715	-2.0	42,700
C44	1036	485	-11.3	5£,300	1179	1148		-3.3 -4.8	43,400 124,900	1260 1261	1629 1555	713	-3.0	42,800
045	1439 1540	407 250	·5.5 -4.2	£7,300 109,200	1180			-5.2	124,900	1262	1468	717 717	-4.0 -5.0	42,600 42,600
C4E	1576	635	3.7	47,100	1181			-5.7 -6.1	125.100	1263	11413	722	-6.0	42,400 -
046	1089	411.	-10.4	€€,700	1183	1363	224	-6.4	: 125:200 124,700	1264 1265	1340		7.0	42.600
050'' 051	, 549 426	.1040 1 818	-13.2 -31.1	26,900 37,800	1164		182	-5.3	164,400	1266	1263 1182	717 720	-8.0 -9.0	42,600 42,500
052	1563	1365	3.6	16,900	1185 1186			-5.8	162.600	1267	1110	717	-10.0	42,500 42,600
csà	779	1092	-16.8	27,000	1189		214	-6.3 -6.2	164,300 131,800	1268 1269	1055	717	-11.0	42,600
054	1613	620	-3.2 -6.5	** 48,000 70.000	1190		286	-5.2	94,200	1270	999 959	717 715	-12.0 -13.0	42,600
055 056	1380 264	377 663	-6.5 <-35.0	72,000 45,500	1191 1152	686 265	1114	-19.5	2€,200	1271	905	712	-13.0	42,700 42,900
C-58	1261	746	-8.0	41,200	1193	.403	ES3 1292	<-35.0 -32.6	34,700 20,000	1272	857	714	-15.0	42,800
260	393	605	33.3	49,000	1154	344	1275	< 35.0	20,600	1273 1274	810 774	705 711		43,300 ···
Æ1 Œ2	1817	645 746	-0.9 -8.2	46,600 41,200	1195 1196	505 572	1311	-27.6	19,400	1277	737	708	-17.0 -18.0	42,900 43,100
64	1258	792	8.1	39,000	1157	572 639	1293 1502	-24,1 -21,2	20,000	1278	702	711	-19.0	42,900
Æ5	705	534	/ -18:9 L	33,000	1198	637	1402	-21.3	13,000 16,300	1279 1280	671 645	710	-20.0	43,000
	\1181 · . 529	734 658	-9.0 -26.3	, 41,800 - 45,800	1199 1200	£14	1467	-22.1	16,200	1281	617	710 707	-21.0 -22.0	43,000 43,100
68	508	696	-27.4	43,700	1200	637 1095	1431 13 9 4	-21.3 -10.3	15,400	1282	595	704	-23.0	43,300
	1898	604	-0.3	49,100	1202	1719	1545	-2.1	16,600 11,600	1283 1284	573 552	700	-24.0	43,500
71 73	£73 1768	609 1128	-14.7 -1.5	48,700 25,800	1203 1204	791 664	668	-16.5	45,200	1285	536	695 694	-25.0 -26.0	43,700 43,800
75	636	773 ,	-15.4	35,900	1205	964 ⊬813	1021 195	-12.9 < 35.0	29,700	1286	515	687	-27.0	44,200
76" . 78	1863	861	-0.6	36,000	1208	306	194	<-35.0	148,700 149,800	1287 1288	496 467	683 669	-28.0	44,400
/6 61	826 571	566 483	-15.7 -12.7	51,600 58,500	1209 - 1210	320	197	<-35.0	147,400	1289	447	667	-29.0 -30.9	45,200 45,300
63	1667	202	-2.3	142,300	1211	326 354	. 197 294	<-35.0 -33.2		1290	. 427.	655	-31.0	45,900
65 90	1157	794	-9.4	3E,900	1212	402	294	32.7	91,400 91,200	1291 1292	412 397	655 652	-32.0	45,900
	620 1867	910 597	-21.9 -0.5	34,000 49,500	1214 1215	386 641	294	-33.7	91,400	1293	381	654	-33.0 -34.0	46,100 ¹¹ 46,000
3	2019	894	>0.0	34,600	1215	660	329 329	-21.2 -20.4	81,600 81,600	1294	365	653	-35.0	46,100
	1546	538	-4.1	53,700	1217	914	266	-13.8	101,800	1295	348	653	<-35.0	46.100
95 98	1545 ' ' 61	477 935	-4.1 35.0	59,100 33,000	1218	873	245	-14.7	112,000					
9	1954	237	>0.0	116,000	1219 1220	970 1021	372 298	-12.7 -11.6	72,900		•			
1		1048	23.3	28,600	1221	1392	205	-6.3	90,100 139,500					
3	1050 457	667 797	-11.1 -29.5	45,200 38,800	1222	1354	203	-6.8	141,800					
5	1884	532	-0.4	54,200	1223 1224	1362 673	205 540	-6.7 -19.9	139,500					
	1714	649	-21	46,300	1225	614	542	-19.9 -22.1	53,600 53,400					
	1717 1976	546 722	-2.1 >0.0	53,100 42,400	1226	603	539	-22.6	53,600					
1		1066	·25.3	28,000	1227 1228	696 707	623 628	-19,2 -18 G	47,800 47,500					
	1348	621	-6.9	48,000	1229	475	447	-18. 9 -28.7	47,500 62,300					
	1385 1078	762 816	-6.4 -10.6	40,400	1230	466	1282	-29.0	20,400					
	975	787	-10.6 -12.6	38,000 39,300	1231 1232	759 1324	1461	-17.4	14,400					
B 1	202	933	-8.7	33,100	1233	1583	1170 1005	-7.2 -3.6	24,200 30,300					
		076	-11.6	27,600	1234	1865	809	-0.6	38,200					
	905 512 1	616 301	-0.3 -4.5	48,300 19,700	1235 1236	1812	817	-1.0	37,900					
2 1	114	677	-9.9	44,700	1237	1411 1392	703 682	-6.0 -6.3	43,400 44,500					
	464	452	-5.1	61,700	1238	794	410	-16.4	66,900					
		857 802	-11.1 -9.8	36,200 38,600	1239	769	407	-17.1	67,300					
1	•	~~~	- J .0	38,600	1240	740	406	-17.9	67,500					

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POP name	Protein name	MSN's	Boole for identification
IDS:3 ALPHA HIDDH			Dasis for longification
	S-a-nyaroxysteroid-dihydrodiol- dehydrogenase, an enzyme of	137, 159	Pure protoin and antibody provided by Dr. T.M.
IDS:ACTIN_BETA	Steroid metabolism B cellular action a conscious		Penning, Department of Pharmacology, School
IDS-ACTIN GAMMA		38	Homologous position with respect to other mammalian
AMMAD	Y cellular actin, a cytoskeletal protein	69	Systems Systems
IDS:ALBUMIN	Serum afbumin, mature form.	21 26 22	Systems
£ - 5	Apo A-I plasma lipoprotein, mature form	236, 463	Predominance in rat plasma
IDS:CALMODULIN	Calmodulin, an acklic cytosofic calcium.	, , , , , , , , , , , , , , , , , , ,	r resence in rat plasma, regulation by some lipid- lowering drugs
IDS-CATALAGE	binding protein	163, 049	Homolognus position with respect to other mammelles
	Catalase (peroxisomal)	54, 61, 106	Present
IDS:CPKSPOTS	Spots contributed by the CPK charge	1257 - 1205	to mouse catalase
iDS:CPS	standards (not rat liver proteins) Carbamoyl phosphate synthase	114, 157, 167, 174, 1184, 1185, 1186, 1222	Pure pro
IDS:CYTOCHROME_B5	Cytochrome b5	87, 477	
IDS:FABP-L	Liver fatty-acid binding protein		Department of Pharmacology, Toxicology and Therapsutics, University of Kansas Medical Conter
	menos Amouno por	777	Pure protein provided by Dr. Nathan Bass, Department
IDS:HMG-COA_SYNTHASE	Cylosolic HMG-CoA Synthase	133, 144, 235, 413	Marting of Translater California School of Milhody provided by Dr. Michael Greenman Mark
IDS:LAMIN_B	Lamin B, a nuclear protein	16 724	Sharp & Dohme Research Laboratories, Rahway, NJ
DS:MITCON:			Homologinus position with respect to other mammallan
C.NOO.	Milcon: 1 (F1 ATPase (I subunit), a	17, 49, 71, 340, 1245, 1246, 1247, 1249	Homohoous position with respect to the contract of the contrac
IDS:MITCON:2	Milcon: 2, a mitochondrial matrix stress	15, 25, 110, 1241, 1242, 1243, 1244	systems, presence in mitochondria
IDS:MITCON:3	protein equivalent to E. Mitcon: 3. a mitochondrial matrix etress	1131 (0131 (1131 (1131 013 3CC 3C 0)	nomologous position with respect to other mammallan Systems, presence in milochondria
De-Mangu para nan	protein, likely analog of	19, 33, 429, 600, 1236, 1239, 1240	Homologous position with respect to other mammallan
IOS.NAUTH_P450_MEU	NAUPH cytochrome P-450 reductase, Irequently co-induced with P-450's	175, 251, 812	Systems, presence in mitochondria Pure profein provided by Dr. Andrew Parkinson, Department of Pharmacology, Toxicology and Thatanadise Inkaperia,
IDS:PDI	Protein disulphide isomerase 1	168, 1170, 1171, 1172	Canter Canter Comment of Canter Cante
IDS:PLASMA_PROTEINS	Rat plasma proteins observed in liver	21, 28, 33, 44, 72, 102, 115, 197, 236, 246, 248, 257, 293, 332, 347, 364, 369, 419, 432, 463, 469, 518, 562, 605, 623, 666, 687, 725	Sequence mormation obtained by R.M. Van Frank, Lilly Research Leboratories, Indianapolis Plasma coelectrophoresis studies
IDS:PRO-ALBUMIN	Serum albumin precursor	738, 790, 865, 903, 926 47, 93	-
IDS:PYRCARBOX IDS:SOD	Pyruvale carboxylase Superoxide dismulase	179, 1180, 1181, 1182, 1183	Pavlica, R.J., et al., 88A (1890) 1022 115-125.
IDS:TUBULIN_ALPHA	a tubulin, a cytoskeletal protein	56, 132, 1224, 1252	Sequence information obtained by R.W. Van Frank, Lily Research Laboratories, Indianapolis
IDS:TUBULIN_BETA	B lubulin, a cytoskalatal protain	50, 1225, 1226, 1251	From the state of the second to other mammallan systems Homologous position with respect to other mammallan

Table 3. Computed pl's of two sets of carbamylated protein standards: Rabbit muscle CPK and human hemoglobin (Hb)

	Protein Name	PIR Name	#ASP 3.9	#GLU 4.1		#LYS 10.8	#ARG 12.5	NH2- 7.0	Calc	Real	er eren gradiag Theodologradiag
-3 -4 -5 -6 -7 -8 -9 -10	Rabbit muscle CPK	_{L.,} KIRBCM	28 28 28 28 28 28	27 27 27 27 27 27 27 27	17 17 17 17 17 17 17 17 17	31 30 29 28 27 26 25 24 23	18 18 18 18 18 18 18 18 18	(a) (q) (b) (q) (b) (q) (a) (a) (a) (a) (a) (a) (a) (a) (a) (a	6.67 6.54 6.42 6.31 6.12 6.03 5.94 5.85 5.76 5.67	2 2 3 1 2 3 1 5 5 5 5 6 5 7 6 5 7 6 5 7 6 9 5 9 9	In of Assente 1883 par environmente 1886 parente per elemente per elem
-13 -14 -15 -16 -17 -18 -19 -20 -21 -22 -23 -24 -25 -26 -27 -28			28 28 28 28 28 28 28 28 28 28 28 28 28 2	27 27 27 27 27 27 27 27 27 27 27 27 27 2	17 17 17 17 17 17 17 17 17 17 17 17 17 1	19 18 17 16 15 14 13 12 11 10 9 8	18 18 18 18 18 18 18 18 18 18	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	5.58 5.48 5.39 5.29 5.20 5.12 5.04 4.96 4.89 4.83 4.77 4.71 4.66 4.61 4.56 4.52 4.48	-12 -13 -14 -15 -16 -17 -18 -19 -20 -21 -22 -23 -24 -25 -26 -27 -28	official Catholighteresis of greather claims of the uplants of the
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5 5 5 7 7 8 9 9			7 7 7 7 7 7 7 7	8 8 8 8 8 8 8 8 8	999999999	8 7 6 5 4 3 2 1 0 0	3 3 3 3 3 3 3 3 3	1 6 1 5 1 5 1 5 1 5 1 4 1 4	5.32 5.13 5.96 5.78 5.59 5.37 5.14 5.91	-3.2 -5.3 -7.2 -10.0 -12.3 -15.5 -18.0 -21.0 -25.5 -27.2	

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Table 4. (Computed of some known proteins related to measured CPK:pfs is a constant of the latest with the formation of the computed of the constant of the co

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O Creatine phospho kinase (CPK), rabbit muscle	JEKIRECM J. D. 28 CC27 Em17 ep/34 (1918) A 6:84 E 5:0.0
Fatty scid-binding protein, ratchepatic Common line	FZATLS 6 - 5 10 13 - 12 - 116 - 2 - 7 83 1 12 0
2 b2-microglobulin, human	MGHUE2 43177 6.8 43.4 4415 8 // 15 6.09 11 -5.0
3 Carbamoyl-phosphate synthase, rather to prove	SYRTCA 72 96 28 28 55 56 5.97 5.5.5
4 Proalbumin (serum albumin precursor), rat	en ABRTS . <32 . 57 - 15 : 53 - 27 . 5.98 - 6.2
5 Serum albumin, rat	ABRTS 32. 57 15 353 24 5.71 6-9.0
6 Superoxid dismutase (Cu-Zn, SOD), ration is a line	A26810 8 3 11 10 10 10 1 4 5 01 0 0
7 Phospholipase C, phophoinositide-specific (?), rat	A28807 34 : -: 42: : 9. x2: 49 : : 21, 5: 92 : -9.2
8 Type Albumin, human the sugar of the state of the	ABHUS and 3661 sant6 review to page 5.70 days of
9 Apo A-I lipoprotein, rat	A24700 18 24 6 23 12 522 12.7
10 proApo A-I lipoprotein, human	LPHUA161 116 300 30 316 3 31 317 5 25 314 0
11 NADPH cytochrome P-450 reductase, rat	hDhilO4 41 60 21 20 20 507 100
12 Retinol binding protein, human and the second	. 35 VAHU- 3 318/5 210 . m .2610 33 314 35:5.04 36-16.9
13 , Actin beta, rat	ATRIC 99 23 9 26 9 9 16 19 13 18 25 5.06 (2) 17.2
14 Actin gamma, rat	ATRTC 20 1 29 . 4.9 4 19 4 18 45.07 416.8
15 Apo A-I lipoprotein, human	LPHUATA
16 Apo A-IV lipoprotein, human	LPHUA4 20 - 49 8 8 285 24 4 4 88 - 19.7
17 Tubulin alpha, rat	UBRTA, 12 27: 3.37 31 13 3 19 30 21 14.66 7 19.8
18 F1ATPase beta, bovine	FWBOB 25 36 9 22 22 4.80 21.0
19 Tubulin beta, pig	* UEPGB 25, 36 or10', 15 22 449* -225
20 Protein disulphide isomerase (PDI), rat hepatic	ISRTSS 43 51 11 51 9 4.07 -25.0
21 Cytochrome b5, rat	CBRT5 10 15 6 10 4 4.59 -26.0
22 Apo C-II lipoprotein, human	LPHUC2 4 7 0 6 1 4.44 -30.5
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High Specific Activity Chemiluminescent and Fluorescent Markers: their Potential Application to High Sensitivity and 'Multi-analyte' Immunoassays

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The sensitivities of immunoassays relying on conventional radioisotopic labels (i.e. radioimmunoassay (RIA) and immunoradiometric assay (IRMA)) permit the measurement of analyte concentrations above ca 107 molecules/ml. This limitation primarily derives, in the case of 'competitive' or 'limited reagent' assays, from the 'manipulation errors arising in the system combined with the physicochemical characteristics of the particular antibody used; however, in the case of 'non-competitive' systems, the specific activity of the label may play a more important constraining role. It is theoretically demonstrable that the development of assay techniques yielding detection limits significantly lower than 107 molecules/ml depends

- (1) the adoption of 'non-competitive' assays designs;
- (2) the use of labels of higher specific activity than radioisotopes;
- (3) highly efficient discrimination between the products of the immunological reactions involved.

Chemiluminescent and fluorescent substances are capable of yielding higher specific activities than commonly used radioisotopes when used as direct reagent labels in this context, and both thus provide a basis for the development of 'ultra-sensitive', non-competitive, immunoassay methodologies. Enzymes catalysing chemiluminescent reactions or yielding fluorescent reaction products can likewise be used as labels yielding high effective specific activities and hence enhanced assay sensitivities.

A particular advantage of fluorescent labels (albeit one not necessarily confined to them) lies in the possibility they offer of revealing immunological reactions localized in 'microspots' distributed on an inert solid support. This opens the way to the development of an entirely new generation of 'ambient analyte' microspot immunoassays permitting the simultaneous measurement of tens or even hundreds of different analytes in the same small sample, using (for example) laser scanning techniques. Early experience suggests that microspot assays with sensitivities surpassing that of isotopically based methodologies can readily be developed.

Keywords: Ultrasensitive immunoassay; fluorescent microspot immunoassay; confocal microscopy

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INTRODUCTION

Immunoassay methods relying on radioisotopic labels have played a major role in medicine and other biologically related fields (agriculture, veterinary science, the food and pharmaceutical industries, etc.) during the past two decades. Their importance has derived from the exploitation both of the 'structural specificity' characterizing antibody-antigen reactions and the 'detectability' of isotopically-labelled reagents, the latter permitting observation of the binding reactions between exceedingly small concentrations of the key reactants involved. The combination of these features endowed radioimmunoassay has use throughout modern medicine and biology. However, in the past few years, interest has increasingly focused on so-called 'alternative', non-radioisotopic, immunoassay methods; such techniques are based on essentially identical analytical principles but differ in the markers used to label the particular immunoreactant (antibody or analyte) whose distribution between bound and free moieties (following the basic analytical reaction) constitutes the assay 'response'. The reasons for this interest may be grouped under four headings:

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- (1) Environmental; logistic; economic; practicality and convenience, etc. (i.e. 'non-scientific).
- (2) The attainment of higher sensitivity.
- (3) The development of 'immunosensors' and 'immunoprobes'.
- (4) The development of 'multi-analyte' assay systems.

Our own reasons for developing non-isotopic techniques fall principally under headings (2) and (4), and this presentation will centre primarily on the concepts which underlie our immunoassay development strategy in these areas.

THE ATTAINMENT OF 'ULTRA-HIGH' IMMUNOASSAY SENSITIVITY

Though, as indicated above, the sensitivity of radioisotopically based immunoassay methods has constituted one of the principal foundations of their widespread use over the past 25 years, a

fundamental reason for their replacement stems, paradoxically, from the current requirement to develop microanalytical techniques which care which superior to them in this particular respect. Radioisotopic methods are, in practice, limited to the measurement of analyte concentrations above about 108-109 molecules/ml (i.e., approx 0.15-1.5) sources pmol/l)(Dakubu et al., 1984). However, in certain fields (e.g. virology, tumour detection) there is a particular need to detect or measure molecular concentrations below this level. The factors which determine immunoassay sensitivity have been extensively discussed (Ekins et al., 1968, 1970a; Ekins, 1978; Jackson et al., 1983; Dakubu et al., methods with unique specificity and sensitivity underlying concepts are still frequently misunder-1984; Ekins, 1985). Nevertheless, some of the stood and merit brief discussion in the present context. The thic bests

The concept of sensitivity

One major source of past confusion has been disagreement regarding the concept of 'sensitivity' itself, many authors equating assay sensitivity. with the slope of the dose-response curve (Yalow and Berson, 1970a, b; Berson and Yalow, 1973; see also Ekins et al., 1970b, Tait, 1970). It is now widely agreed that the notion that a steeper dose-response curve implies greater sensitivity is erroneous. The invalidity of this belief is clearly revealed by the fact that the relative magnitudes of the responses yielded by two assay systems is dependent on the particular variable which is chosen to represent the response (see Fig. 1(a))(Ekins, 1976). For this and other reasons, it has long been recognized that the 'sensitivity' of an assay can only be satisfactorily represented by its lower limit of detection (Fig. 1(h)), and this concept is now embodied in all internationally agreed definitions of the term. An essentially identical definition is as the precision (i.e. standard deviation) of measurement of zero dose, since this quantity determines the least quantity distinguishable from zero and hence the assay detection limit. The sensitivity of an assay is thus represented by the zero-dose intercept of the 'precision profile' (Fig. 2(a)) when the latter is expressed in terms of standard deviation rather than of coefficient of variation (Ekins, 1983a). In short, the more sensitive of two assays is the one yielding greater precision of the zero dose estimate (Fig. 2(b)).

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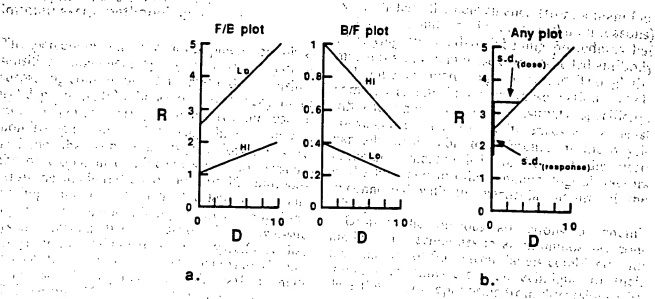


Figure 1. (a) Diagrammatic representation of conventional RIA dose-response curves for systems using high (hi) and low (lo) antibody concentrations plotted in terms of free-bound (F/B) and bound/free (B/F) labelled antigen. Note that the use of a lower amount of antibody yields a dose-response curve of greater slope in the F/B plot, but of lower slope in the B/F plot. It is impossible to decide, on the basis of the data shown in this figure, which concentration of antibody yields the assay system of higher sensitivity. (b) The sensitivity of an assay is essentially represented by the minimum detectable dose, i.e. the SD of the dose measurement (SD_{toose)}) at zero dose. This is given by the SD of the response (SD_{tresponse}) divided by the dose-response curve slope at zero dose (i.e. $((SD_{tresponse})) \times dD/dR)_0$). This quantity is unaffected by the choice of the coordinate frame used to plot the dose-response curve. (Note: it is common to multiply (SD_(gose))₀ by an arbitrary factor to increase the confidence level ettaching to the minimum detectable dose estimate, though, since no agreement exists regarding the value of this factor, this unnecessary step merely adds to confusion when the relative sensitivities of two assay procedures are compared.)

'Competitive' and 'non-competitive' ('limited reagent' and 'excess reagent') assays

A second important misconception in this area is the notion that immunoassays relying on the use of labelled antibodies (e.g. immunoradiometric assays, IRMA) are ipso facto more sensitive than

those which rely on the use of labelled 'analyte' (e.g. radioimmunoassays, RIA); furthermore the grounds originally advanced for the claimed superiority of labelled antibody methods (Miles and Hales, 1968) were partially based on false concepts of sensitivity, and thus failed to identify the true reasons why certain assay designs are

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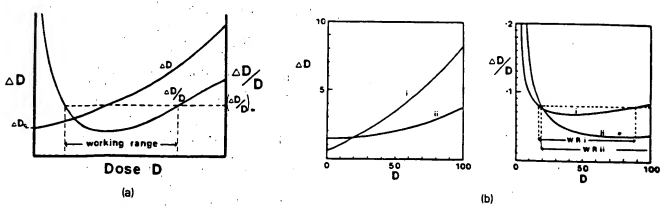


Figure 2. (a) The 'precision profile' of an assay portrays the error in the dose measurement as a function of dose. The error may be represented, inter alia, by the absolute error (ΔD ; e.g. SD of D) or the relative error ($\Delta D/D$; e.g. CV of D). (ΔD)₀, the error in the measurement of zero dose, represents the sensitivity of the assay. The working range may be defined as the range of dose values within which $\Delta D/D$ is less than an 'acceptable' value set by the investigator. (b) The more sensitive of the two assays (assay I) intercepts the ΔD axis at a lower value. However, assay II is more precise at higher values of dose, and has a wider working range.

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potentially capable of yielding far higher sensitivity than others. This issue likewise merits clarification:

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The purely pragmatic sub-classification of immunoassays into labelled antibody and labelled analyte methods diverts attention from a more fundamental divide in immunoassay methodology, which relates to the optimal concentration of antibody required in an assay system to maximize its sensitivity. In certain assay designs (which may be termed 'limited reagent' or 'competitive') the optimal concentration tends to zero; conversely in others (which may be termed 'excess reagent' or 'non-competitive') the concentration tends to infinity. It should be particularly emphasized that the optimal antibody concentration is essentially governed, not only by the physicochemical characteristics of the antibody-analyte binding reaction, but also by the errors incurred in measurement of the assay response. Were an assay system to be totally error-free, no antibody concentration would be optimal, and the distinction between competitive and non-competitive methodologies would thus not arise.

Though it is inappropriate in this presentation to discuss in detail the statistical and physicochemical theory underlying this fundamental divergence in immunoassay design (see Ekins et al., 1968, 1970a; Jackson et al., 1983), the reason for it can perhaps be more readily understood if the basic principles of immunoassay are portrayed in a somewhat different way from that in which they are usually presented. All immunoassays essentially depend upon measurement of the 'fractional occupancy' by analyte of antibody binding sites following reaction of analyte with antibody (see Fig. 3(a)). Those techniques which implicitly rely on measurement of residual, unoccupied, binding sites optimally necessitate the use of concentrations of antibody tending to zero, and may be termed 'competitive', conversely those in which occupied sites are directly measured necessitate use of high antibody concentrations and are termed 'non-competitive' (Fig. 3(b)). This emphasizes that the differences in assay design characterizing so-called competitive and non-competitive methods are essentially unrelated to which component (if any) of the reaction system is labelled. Indeed immunoassays in which no label of any kind is involved can, on identical grounds, be subdivided into those of 'limited reagent' (or 'competitive') and 'excess reagent' (or 'non-competititve') design. Thus the

distinction between these two forms of immunoassay simply reflects differences in the way that fractional antibody occupancy is determined, and the fact that it is generally undesirable—for reasons of accuracy—to measure a small quantity by estimating the difference between two large quantities. When an immunoassay relies on the measurement of unoccupied antibody binding sites, the total amount of antibody used in the system must be small to minimize error in the resulting (indirect) estimate of occupied sites.

Measurement of occupied sites

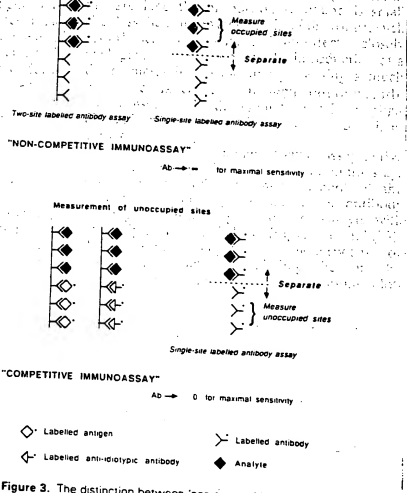


Figure 3. The distinction between 'non-competitive' (above) and 'competitive' immunoassays (below) reflects how antibody binding-site occupancy is measured. Labelled antibody methods are 'non-competitive' if occupied sites of the (labelled) antibody are measured, but are 'competitive' (below right) when unoccupied sites are measured. Labelled antigen (below left) or labelled anti-idiotypic antibody methods (below centre) rely on measurement of sites unoccupied by analyte, and are therefore invariably, of 'competitive' design.

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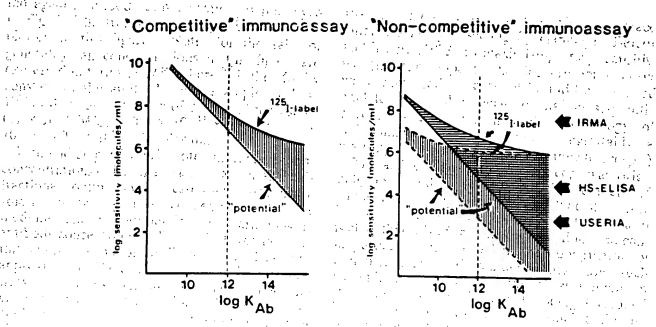


Figure 4. Curves showing the theoretically predicted relationship between antibody affinity and the sensitivities achievable using 'competitive' and 'non-competitive' assay strategies. The 'potential' sensitivity curves assume the use of infinite specific activity labels; the sensitivities achievable using 1251-labelled antigen or antibody are also shown. Shaded areas indicate the sensitivity loss due to errors in measurement of the label. Curves relating to 'competitive' assays assume a 1% error in measurement of the response variable arising from 'experimental' errors (i.e. errors other than those inherent in label measurement per se). Non-competitive curves assume 'non-specific binding' of labelled antibody of 0.01% and 1% (lower and upper curves) respectively. Arrows indicate sensitivities claimed for typical non-competitive immunoassay methodologies.

Conversely, when occupied sites are measured directly, this particular constraint does not arise; indeed, considerable advantage often derives from using relatively large amounts of antibody in the system.

Sensitivity of 'competitive' and 'non-competitive' immunoassays

Competitive and non-competitive immunoassays differ significantly in many of their performance characteristics in consequence of the differences in optimal antibody concentration on which they rely. Most particularly they differ in their potential sensitivities. Figure 4. portrays the sensitivities predicted theoretically as a function of antibody binding affinity, making realistic assumptions regarding the experimental errors incurred in reagent manipulation, 'non-specific' binding of labelled antibody, etc., and assuming the use of optimal reagent concentrations (Ekins, 1985). Amongst other concepts illustrated in the figure is the much greater assay sensitivity potentially attainable (using an antibody of given affinity) by adoption of a non-competitive approach. In short, whereas the maximal sensitiv-

ity realistically achievable using a competitive design is in the order of 10⁷ molecules/ml (using antibody of the highest affinity found in practice), a non-competitive method is capable of yielding sensitivities some orders of magnitude greater than this. However, Fig. 4 also demonstrates that. assuming the use of high affinity antibodies (i.e. $\sim 10^{11} - 10^{12} \, \text{l/m}$), maximal sensitivities yielded by isotopically based techniques (whether relying on labelled antibody (IRMA) or labelled analyte (R1A), or whether of competitive or noncompetitive design) are closely comparable, i.e. of the order of 10^7-10^8 molecules/ml.

This limitation is a manifestation of the fact that, in the case of the non-competitive methods. an important constraint on assay sensitivity is (under certain circumstances) the 'specific activity' of the label used. On the other hand, limitation of assay sensitivity due to the low specific activity of radioisotopic labels does not often arise, in practice, in the case of competitive assays, whose sensitivity is generally restricted by other factors (Ekins, 1985). The fundamental significance of this conclusion is that, only by the use of labels possessing specific activities higher than those of the commonly used radioisotopes in assays of non-competitive design, can current

THE MITCHAEL SECRET OF FUTURES CER sensitivity limits be breached. Conversely, use of a higher specific activity label in a competitive assay will usually have no significant effect on its sensitivity (assuming experimental errors incurred in reagent manipulation of the magnitude generally encountered in practice). antient and months of the first state of the second of t

High specific activity non-isotopic labels

the and the stage of the entropy of the The term 'specific activity' is conventionally applied, in the case of radioisotopic labels, to denote the number of radioactive disintegrations per unit time per unit weight of the isotope or labelled compound. In the present context, use of the term is widened to signify 'detectable events' per unit time per unit weight of labelled material. Thus it can be used to indicate the rate of photon emission by a chemiluminescent or fluorescent label, or the rate of conversion of substrate molecules-by an enzyme label-to molecules of a detectable product. The importance of the concept derives from the fact that 'signal measurement error' (i.e. error in the measurement of the label per se) is a contributory factor in limiting assay sensitivity, and may-when other sensitivity-constraining factors are reducedbecome dominant. Furthermore, when extending the sensitivities of immunoassay systems beyond their present limits, the numbers of molecules involved are low, and statistical errors incurred in counting individual 'detectable events', and the time required to count them, may assume a particular importance.

Table 1 compares the specific activities of potentially useful labels with that of 1251. All are of relevance in the context of this volume since chemiluminescent and fluorescent labels can be used to label antibodies (or antigens) directly; alternatively, enzyme labels catalysing reactions yielding chemiluminescent signals or fluorescent products can be utilized.

The importance of background in non-competitive immunoassays

A second important factor governing the sensitivity of non-competitive labelled-antibody immunoassays is the 'background' or 'blank' signal emitted in the absence of analyte, since error in the measurement of this signal is clearly a major determinant of the error in measurement of zero

Table 1. Relative specific activities of various isotopic and non-isotopic labels. Note that, though the specific activity of 1251-labelled reagents does not, in practice, significantly limit the sensitivity of the procompetitive assays (see Fig. 4), the lower specific activity of ³H may severely restrict the sensitivity of competitive assays (e.g. of steroid hormones) which rely on the use of this particular radioisothe action, participated and The rest of the wife the

Specific Activities

detectable event/sec/7.5 x 106 so labelled molecules: 4 a it will be a 1 detectable event/sec/5.6 x 10⁸ labelled molecules. Enzymes: Determined by enzyme 'amplification factor and detectability of reaction product. Chemiluminescent 1 detectable event/labelled molelabels cuie. Fluorescent labels: Many detectable events/labelled molecule.

dose. Amongst contributors to the background signal are the 'noise' of the measuring instrument itself, 'ambient' signal generators (such as, inimmunoassays, solid 'captureantibody' supports or, in the case of radioisotopic methods, cosmic ray and other extraneous radiation sources) and 'non-specifically bound' labelled antibody. Minimization of each of these components is essential for maximal sensitivity: mere arithmetic subtraction of background is of absolutely no benefit in this context.

Non-specific binding of antibody is of particular interest, since the magnitude of this contribution is dependent, inter alia, on the amount of labelled antibody used in the system, and the duration of its exposure to analyte. Thus increasing the amount of labelled antibody increases the amount of such antibody bound to analyte; however, it may also increase the non-specifically bound moiety to a greater proportional extent, and thus cause a net reduction in sensitivity. This effect underlies the loss in sensitivity at higher antibody concentrations depicted in Fig. 5 (reproduced from Jackson et al., 1983). This phenomenon also underlies the relationship between sensitivity and the affinity constant of the labelled antibody depicted in Fig. 4. The possession by labelled antibody of a high affinity constant implies that a

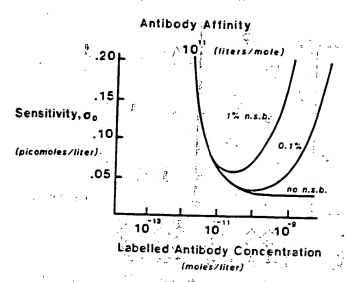


Figure 5. Assay sensitivity (represented by the standard deviation of the zero dose measurement, o₀), plotted as a function of the concentration of labelled antibody (of affinity 10¹¹ L/M) used in the assay, assuming different levels of non-specific binding of labelled antibody. (Note: an irreducible instrument background has been assumed in the computations represented; this limits the ultimate sensitivity attainable, regardless of the concentration of antibody used.)

lower concentration is required to yield the same level of analyte binding, albeit with reduced non-specific binding, thus increasing assay sensitivity

In summary, the high sensitivity of noncompetitive labelled antibody methods derives essentially from their permitted use of optimal concentrations of antibody which (provided nonspecific binding of labelled antibody is low) are generally considerably greater than in competitive methods, not from the fact that the antibody is labelled. Labelled antibody methods generally fall in sensitivity as the concentration of antibody is reduced towards zero, ultimately yielding a sensitivity theoretically identical to that of competitive methods (Rodbard and Weiss, 1973). (Paradoxically, early exponents of labelled antibody methods, whilst claiming them to be of higher sensitivity, also concluded that their sensitivity was increased by reduction in the amount of labelled antibody used (Woodhead et al., 1971). This incorrect conclusion—based on observation of effects on the slope of the dose-response curve—exemplifies the many fallacies encountered in the immunoassay field stemming from confusion regarding the concept of sensitivity discussed above.) Finally it should be

emphasized that maximization of the sensitivity of a non-competitive immunoassay generally implies the selection of reagent concentrations and other experimental conditions such that the [analyte signal/background] ratio (i.e. s/b) is maximized. However, this simple relationship disregards statistical considerations which arise when the numbers of detectable events are very low, and a more appropriate objective may, under these circumstances, be maximization of the ratio s^2/b (Loevinger and Berman, 1951).

Other performance characteristics of competitive and non-competitive immunoassays

Non-competitive designs also display a number of other advantages deriving from the relatively high antibody concentrations on which they generally rely. These include increased reaction speeds (and hence shorter incubation times), decreased vulnerability to certain environmental effects (which cause variations in binding affinity between antibody and analyte), reduced sensitivity-dependence on high antibody binding affinity, etc.

Nevertheless a price has to be paid for these benefits; this includes the greater tendency of a large amount of antibody to bind molecules differing from, but with structural resemblance to, the analyte itself, implying a loss of assay specificity. This effect generally necessitates the use, whenever possible, of an 'immunoextraction' procedure using a second 'capture' antibody (usually directed against a different binding site, or 'epitope') as shown in Fig. 3(b). This technique-the 'sandwich' or 'two-site' immunoassay (Wide, 1971)—thus potentially combines the twin virtues of ultra-high sensitivity and specificity (together with short reaction time), features of crucial importance in many diagnostic situations (for example, in the detection of AIDS viral antigens). (Note, however, that the loss of specificity inherent in non-competitive assay designs implies that they are less readily applicable to the measurement of analytes of small molecular size, which cannot be simultaneously bound by two different antibodies directed against different antigenic sites on the molecule. Such analytes are generally more appropriately measured using 'competitive' assay methods.)

1.33 Development of ultra-sensitive immunoassay methodologies

The perception that the development of fultrasensitive' immunoassay systems (i.e. systems surpassing conventional RIA methods in sensitivity) depends on (a) reliance on 'excess reagent' or activities than commonly used radioisotopes; (c) the development of efficient separation systems (ensuring minimization of non-specific antibody binding, and hence of signal 'backgrounds'), and (d) dual or multi-antibody analyte-recognition systems (exemplified by 'sandwich' or two-site assays) to maintain/increase assay specificity, has formed the basis of our own laboratory's immunoassay development since the early to mid-1970s (Ekins, 1978). This led us, inter alia, to an immediate recognition (Ekins, 1979, 1980) of the importance of the in vitro techniques of monoclonal antibody production pioneered by Köhler and Milstein (1975), which are currently the subject of bitter patent disputes in the USA (Ezzell, 1986, 1987a,b), and which may be expected in Europe.

Meanwhile, of the candidate labels for use in this context, both chemiluminescent and fluorescent labels offer many attractions. The development of stable, highly chemiluminescent, acridinium esters by McCapra and his colleagues (McCapra et al., 1977) has subsequently been exploited by Weeks et al (1983, 1984) and, more recently, by several commercial kit manufacturers; other workers have used more conventional chemiluminescent compounds to label immunoassay reagents (see, for example, Kohen et al., 1984, 1985; Barnard et al., 1985). Yet others have relied on enzyme labels to catalyse chemiluminogenic (Whitehead er al., 1983) and fluorogenic (Shalev et al., 1980) reactions as indicated above. Detailed description of these various methodologies is presented by others in this volume and need not be duplicated here.

Common to all the 'ultra-sensitive' immunoassay methodologies relying on such alternative labels is their dependence on a non-competitive, labelled antibody, assay strategy whenever appropriate; however, for the reasons indicated above, competitive methods continue to be generally employed for the measurement of analytes of small molecular size (e.g. therapeutic drugs, steroid and thyroid hormones, etc.).

Nevertheless, the convenience (from a manufacturing viewpoint, and for other technical reasons) of relying on standard labelling procedures has meant that, even in these cases, labelled antibody techniques are increasingly preferred. Though the commercial kits based on these various labels 'non-competitive' assay designs; (b) the use of convenience, etc., such differences are at least differ to a minor extent in sensitivity, specificity, non-isotopic labels displaying higher specific partially attributable to differences in the physicochemical characteristics of the antibodies used in the kits, and to other 'immunological' factors unconnected with the particular nature of the label per se.

Despite the obvious attractions of chemiluminescent techniques in an immunoassay context, the use of fluorescent labels combined with sophisticated time-resolution techniques for their detection (a concept arising from discussions with J. F. Tait in 1970) appeared to us (in the mid-1970s) to offer more exciting long-term possibilities for a number of reasons. These naturally included attainment of the enhanced specific activities and high signal to background ratios required for ultra-sensitive immunoassay as indicated above. However, more importantly, fluorescence techniques also appeared to provide a simple route to the development of 'multianalyte' assay systems of the kind described

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In pursuance of this strategy, we began collaboration with LKB/Wallac, ca 1976-77, in the development of the instrumentation and technology required to develop such methods. Fortunately a group of fluorescent substances generally known as the lanthanide chelates (including, in particular, the chelates of europium, samarium and terbium facilitate such development, possessing prolonged fluorescence decay times (~10-1000 µs), large Stokes shift (~300 nm) and other desirable physical characteristics which permit the construction of relatively cheap instrumentation for their measurement (Marshall et al., 1981; Hemmilä et al., 1983). The fluorescent properties of the lanthanide chelates may be compared with those of a conventional fluorophor such as fluorescein which is characterized by a much smaller Stokes shift (~28 nm), and a fluorescent decay time and emission spectrum which imply that it is less readily distinguished from fluorescent substances present in blood (such as bilirubin) or in plastic sample holders. The unique fluorescence characteristics of the lanthanide chelates thus permit them to be

familia wa kasa manaya na gajiya ya ga measured in the presence of a fluorescence background (deriving from extraneous sources) which, in practice, approaches zero. Fig. 6 illustrates the basic concepts involved in pulsedlight, time-resolved, fluorescence measurement, which form the basis of the DELFIA immunoassay system currently marketed by LKB/Wallac.

Though it is inappropriate to pursue this subject in greater detail, attention should also be drawn to the possibilities offered by phaseresolved fluorimetry. This permits separate idenlifetime by their exposure to a sinusoidally modulated exciting light source, and observation. of their demodulated, phase-shifted, light emission (McGown and Bright, 1984). This technique offers the possibility both of the development of homogeneous assays (relying on a difference in fluorescence decay time of bound and free forms. of the fluorescent-labelled molecule), and of discriminating between two labelled antibodies in the context of multi-analyte 'ratiometric' immunoassay as discussed below.

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'AMBIENT ANALYTE' IMMUNOASSAY

Before proceeding to a discussion of the development of multi-analyte assays, another important concept, termed 'ambient analyte immunoassay' (Ekins, 1983b), must first be examined. This term is intended to describe a type of immunoassay system which, unlike unconventional

Intensity Time photon counting time excitation pulse

Figure 6. Basic principles of pulse-light, time resolved fluorescence. Fluorescence emitted by the fluorophor (typically a europium chelate) is distinguished from background fluorescence, which decays more rapidly.

methods, measures the analyte concentration in the medium to which an antibody is exposed, being essentially independent both of sample volume, and of the amount of antibody present. This concept is illustrated in Fig. 7, and relies on the physicochemically-based proposition that, when a 'vanishingly small' amount of antibody (preferably, but not essentially, coupled to a solid support) is exposed to an analyte-containing medium, the resulting (fractional) occupancy of antibody binding sites solely reflects the ambient tification of fluorophores differing in fluorescence analyte concentration. Clearly the binding by antibody of analyte results in a depletion of the amount of analyte in the surrounding medium, but provided the proportion so bound is small (i.e. less than, for example, 1% of the total), such disturbance can be ignored. (This effect is closely analogous to that caused by the introduction of a thermometer into a medium possessing a much larger thermal capacity; the temperature disturbance caused by the thermometer itself is negligible and can, in these circumstances, be disregarded.)

The principles of ambient analyte assay derive from the recognition that all immunoassays essentially depend upon measurement of the 'fractional occupancy' by analyte of antibody binding sites following reaction of analyte with antibody as discussed above (Figs 3. (a) and (b)). The fractional occupancy of ('monospecific' or 'monoclonal') antibody binding sites in the presence of varying analyte concentrations, plot-

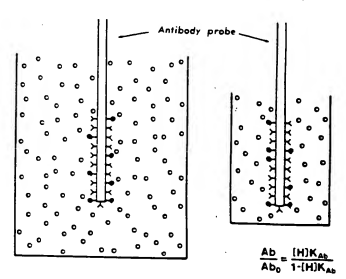


Figure 7. Basic principle of 'ambient analyte' immunoassay (AAI). The fractional occupancy (F) of a vanishingly small amount of antibody (of affinity K) is determined by the analyte concentration in the medium ([An]).

ted against antibody concentration, is portrayed in Fig. 8. The fraction of analyte bound is also plotted in this figure. (Note: for the sake of generality; all concentrations in this figure are expressed in terms of 1/K, where K is the affinity constant of the antibody. For example, if $K = 10^{11} \text{ L/M}$, a concentration of $0.1 \times 1/K$ represents $0.1 \times 10^{-11} \text{ M/L}$, or $0.1 \times 10^{-11} \times 10^{-3} \times 6.02 \times 10^{23} = 6.02 \times 10^8 \text{ molecules/ml.})$

It should be particularly noted that, at antibody concentrations of less than $ca~0.01 \times 1/K$ antibody fractional occupancy is essentially dependent solely on the analyte concentration in the medium, and is independent of variations in antibody concentration. This reflects the fact that this concentration of antibody binds less than approximately 1% of the analyte in the medium, irrespective of its concentration. This implies, for example, that the introduction of 10, 100, or 1000 antibody molecules into a medium containing billions of analyte molecules will result, in each case, in virtually identical fractional antibody binding-site occupancy, the upper limit of antibody concentration being determined by the antibody affinity constant. (An antibody concentration of $0.01 \times 1/K$ is a hundred-fold less than

that (1 × 1/K) necessary to bind 50% of a 'trace' amount of analyte (see Fig. 8), claimed by Berson and Yalow (1973) as maximizing assay 'sensitivity' (i.e. the slope of the dose-response curve when expressed in terms of bound/free labelled analyte). This false conclusion has subsequently become incorporated into the mythology of radioimmunoassay design which, regrettably, a majority of kit manufacturers continue to accept.)

The ambient analyte assay concept was originally exploited in the original development of what has come to be known as 'two-step' free hormone immunoassay (Ekins et al., 1980), but it is clear that it is of far wider application, and can, in particular, be utilized in the construction of immunosensors and immunoprobes. One such example is a probe for the measurement of salivary steroids that is currently being developed in our laboratory. Comprising a small antibodycoated plastic 'dipstick' comparable in size and shape to a clinical thermometer, this device is intended to permit the measurement of salivary steroid levels without requiring the collection of saliva. However, the concept also underlies our approach to multi-analyte immunoassay, also under development in our laboratory.

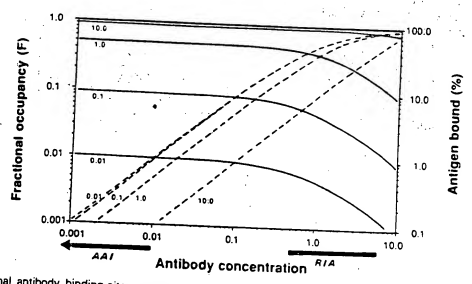


Figure 8. Fractional antibody binding-site occupancy (F) plotted as a function of antibody binding-site concentration for different values of analyte (antigen) concentration [An]. The percentage binding of analyte to antibody (b) is also shown. All becomes binding of analyte is <1%, and fractional binding-site occupancy is essentially unaffected by variations in antibody other competitive immunoassays are commonly designed using antibody concentrations approximately 0.5/K-1/K or above implying $b_0 > 30\%$), in accordance with the precepts of Berson and Yalow (e.g. Berson and Yalow, 1973).

MULTI-ANALYTE 'RATIOMETRIC' **IMMUNOASSAY SYSTEMS**

The concepts relating to ambient analyte immunoassay and assay sensitivity outlined above are both exploited in our present development of a random access, multi-analyte, immunoassay technology capable of measuring, in the same small sample, virtually any number of individual analytes from selected analyte 'menus' (e.g. a hormone menu, viral antigen menu, an allergen menu, etc.). Many examples of a need to measure a multiplicity of different analytes in the same sample exist in medical diagnosis, for example, in the routine diagnosis of thyroid disease, where it different hormones and thyroid-related proteins. At present, clinicians frequently experience difficulty in deciding on the best sequence of tests to arrive at a correct diagnosis. Such problems would be overcome were all relevant analytes measurable at a cost comparable to the cost of measurement of a single substance. Our own immediate objective is the development of a technology permitting the measurement of complete 'hormone profiles' using a single small blood sample. However, the need for 'multi-analyte', or 'random access' measurement is not confined to medical diagnosis: it also arises, for example, in the pharmaceutical industry (where there exists a requirement to ensure the purity of protein drugs synthesized by recombinant DNA techniques), in the food industry and elsewhere. Though still at an early stage, our approach to the achievement of this objective can be briefly indicated.

Multi-analyte assay: general principles

As discussed above, the notion of ambient analyte assay simultaneously introduces two extremely important and novel concepts: (a) that an estimate of analyte concentration can be based upon the use of an infinitesimal amount of 'sampling' antibody, and (b) that such an estimate derives from a direct measurement of fractional antibody occupancy by analyte, irrespective of the exact amount of antibody used. It should be emphasized that the latter proposition is valid only in the context of ambient analyte assay, and is not true in current conventional immunoassay systems (in which fractional antibody occupancy depends both upon the amount of antibody in the

system, and sample volume—see Fig. 8). In short, exposure of a small number of antibody molecules (in the form, for example, of a 'microspot' located on a solid support) to an analytecontaining fluid results in occupancy of antibody binding sites in the microspot reflecting the analyte concentration in the medium. Following such exposure, the antibody-bearing probe may be removed and exposed to a 'developing' solution containing a high concentration of an appropriate second antibody directed against either a second epitope on the analyte molecule if this is large (i.e. the occupied site), or against unoccupied antibody binding sites in the case of small analyte molecules (see Fig. 3(b)). (Note: an is frequently necessary to measure a number of antibody simulating antigen, and reacting with unoccupied binding sites, is described as a 'mirror-image anti-idiotypic antibody'; the use of such an antibody instead of labelled antigen is convenient but not essential, and is suggested here merely to simplify illustration of the basic concepts involved.)

Subsequently, an estimate of binding-site occupancy of the 'sampling' (solid phase) antibody located in the microspot may be derived by measurement of the ratio of signals emitted by the two antibodies forming the dual-antibody 'couplets'. This can be conveniently achieved by labelling the 'sampling' and 'developing' antibodies with different labels, for example, a pair of radioactive, enzyme or chemiluminescent markers. Fluorescent labels are nevertheless particularly useful in this context because, by the use of optical scanning techniques, they permit arrays of different antibody 'microspots' distributed over a surface, each directed against a different analyte, to be individually examined, thus enabling multiple assays to be simultaneously carried out on the same small sample. Fig. 9 illustrates these basic ideas, and Fig. 10 such an array.

Microspot immunoassay sensitivity: theoretical considerations

The notion that it is, in principle, possible to measure an analyte concentration using a microspot of antibody comprising a number of antibody molecules in the range ca 101-106 is likely, at first sight, to appear surprising, and may, indeed, provoke scepticism regarding the assay sensitivities potentially attainable using this approach. Clearly a number of factors, such as the sensitivity

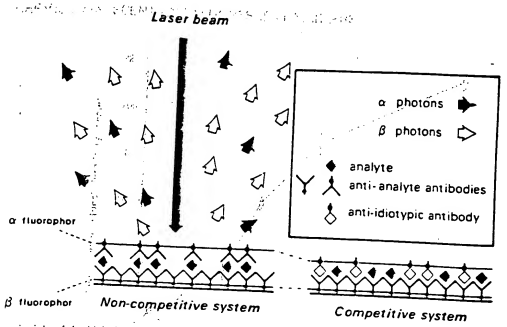


Figure 9. Basic principle of dual-label, ambient-analyte, immunoassay relying on fluorescent labelled antibodies. The ratio of α and β fluorescent photons emitted reflects the value of F (see Figs 5 and 6) and is solely dependent on the analyte concentration to which the probe has been exposed. It is unaffected by the amount or distribution of antibody coated (as a monomolecular layer) on the probe surface.

of the signal measuring equipment, the density of antibody molecules on the surface of the solid support, etc., are likely to play a part in determining final assay sensitivity. Such factors are, in turn, dependent on the efficiency with which the particular labels used can be detected, the adsorption properties of antibody supports,

Figure 10. 'Multi-analyte' antibody array. Each antibody 'microspot' represents a 'vanishingly small' amount of antibody directed against an individual analyte.

able estimates can be made of the order of sensitivities likely to be achieved on the basis of some simple theoretical calculations. To clarify the following discussion, it is assumed that 'sensing' antibody can be uniformly and consistently coated on a solid matrix at a standard density, implying that only the 'developing' antibody need be labelled and measured in order to ascertain fractional occupancy of sensing antibody binding sites.

Fig. 11 illustrates the surface of an antibody microspot, of surface area $A(\mu m^2)$, and (uniformly) coated with antibody of affinity K(L/M) in a monomolecular layer of density D(molecules/μm²). Let us assume that the spot is exposed to an analyte-containing medium of volume v(ml), and containing an analyte concentration C molecules/ ml. The molecular concentration of antibody in the system is thus given by AD/v. (Note: the fact that antibody is situated on the surface of a solid support, and not evenly distributed throughout the medium, does not affect the extent of analyte binding at thermodynamic equilibrium, assuming that antibody binding sites are not impeded in their reactions and have not been damaged during the coating process.)

Meanwhile, fractional occupancy (F) of antibody binding sites by analyte (at equilibrium) is

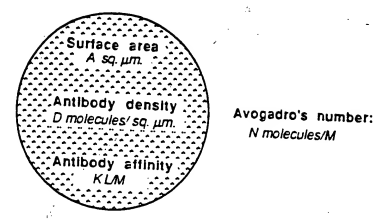


Figure 11. Microspot ambient-analyte immunoassay. The microspot shown is assumed to be uniformly coated with antibody, though if the dual-labelled antibody 'ratiometric' approach shown in Fig. 9 is adopted, uniform coating is not essential. The minimum fluid volume for ambient analyte assay conditions to prevail (enabling adoption of the ratiometric approach) is shown. Minimum test sample volume (M/S): $A \times D \times K \times 10^5/N$

given by the equation:

$$F^2 - F(1/q + p/q + 1) + p/q = 0$$
 (1)

where p = analyte concentration, q = antibody concentration (both expressed in units of 1/K).

Thus, for antibody binding site concentrations $\rightarrow 0$ (i.e. q < 0.01), $F \approx p/(1+p)$; (see Fig. 8). Likewise, the fraction of analyte bound by antibody (f) at equilibrium is given by the equation:

$$f^2 - f(1/p + q/p + 1) + q/p = 0$$
 (2)

Thus, for analyte concentration $\rightarrow 0$ (i.e. p < 0.01), $f \approx q/(1+q)$; (see Fig. 8). Furthermore, when q < 0.01, and when $p \ge 0$, f < 0.01.

Expressed in units of 1/K; the concentration (q) in the assay of 'sensing' antibody situated on the microspot is given by $DAK/(v \times 6 \times 10^{20})$, (since Avogadro's constant, expressed as the number of molecules/mmol, is 6×10^{20} (approximately)). The fraction of an analyte concentration $\rightarrow 0$ which will be bound to the spot is therefore $DAK/(v \times 6 \times 10^{20} + DAK)$, implying that the number of analyte molecules bound to the spot is given by $vCDAK/(v \times 6 \times 10^{20} + DAK)$.

Case 1: sandwich (two-site) assay. Following incubation of sample with antibody, we assume the sample is removed, and the microspot then exposed to a volume V(ml) of a solution of a second, labelled, 'developing' antibody of affinity K^* (L/M) at a concentration given by Q (expressed in units of $1/K^*$).

The fraction of analyte bound by labelled antibody (F^*) at equilibrium is given by the equation:

$$F^{*2} - F^*(1/P + Q/P + 1) + Q/P = 0$$
 (3)

where P represents the analyte concentration in the developing-antibody solution, expressed in units of $1/K^*$, i.e. $\nu CDAKK^*/[(\nu \times 6 \times 10^{20} + DAK)V \times 6 \times 10^{20}]$.

Assuming P < 0.01, $F^* \approx Q/(1 + Q)$. (For example, if Q = 1, the fraction of analyte molecules bound by labelled antibody = 0.5 approximately). Thus, since the number of analyte molecules bound to the spot is given by $\nu CDAK/(\nu \times 6 \times 10^{20} + DAK)$, the number of analyte molecules labelled by the second, developing, antibody is given by $\nu CDAKQ/[(\nu \times 6)]$ $\times 10^{20} + DAK((1 + \tilde{Q}))$, and the surface density of such molecules is given by $vCDKQ/[(v \times 6 \times$ $10^{20} + DAK$) (1 + Q)]. Moreover, assuming that $DAK \ll v \times 6 \times 10^{20}$ (i.e. that the amount of antibody in the system is such that 'ambient assay' conditions prevail, then the surface density $(D^{\tilde{\bullet}})$ of developing-antibody molecules = CDKQ/[(6 $\times 10^{20}$)(1 + Q)] approximately. It should be noted that D^* is independent of both ν and V, also that the ratio $D^*/D = C \times KQ/[(6 \times 10^{20})(1$ + Q)] = $C \times constant$.

If the minimum detectable surface density of developing-antibody molecules (i.e. σ_{D0} , the standard deviation of the measurement of D^{\bullet} when C=0) is given by D_{\min}^{\bullet} (molecules/ μ m²) and C_{\min} represents the minimum detectable analyte concentration in the test sample, then,

disregarding non-specific binding of developing antibody within the microspot area, and the

1.25%,我们并介绍在30%,是严肃的特别,我也们的"是被告诉"。 1.35%,我们们的"是我们"

$$C_{\min} = D_{\min}^* \times [(6 \times 10^{20})(1 + Q)]/DKQ$$
 (4)

 $C_{\min} = D_{\min}^* \times [(6 \times 10^{20})(1 + Q)]/DKQ$ (4) For example, if Q = 1, $D = 10^5$ molecules/ μ m², K= 10^{11} L/M and D_{\min}^* = 20 molecules/ μ m², then $C_{\text{min}} = 2.4 \times 10^6 \text{ molecules/ml} = 10^{-15} \text{ M/L}. \text{Jt}$ should be noted, in this example, the fractional occupancy of the sensing antibody binding sites. by the minimum detectable analyte concentration... is 0.04%.

Case 2: anti-idiotypic antibody ('competitive') assay. In this case, we assume that, following removal of the sample, the microspot is exposed to a volume V(ml) of a solution of (for example) a second, labelled, anti-idiotypic antibody reacting with unoccupied sites on the sensing antibody. Using similar reasoning as above, we may likewise assume that the fraction of such sites which become occupied by the anti-idiotypic developing' antibody is given by Q/(1 + Q), where Q is the developing-antibody concentraion. However, the minimum detectable surface density of anti-idiotypic antibody is not, in a competitive design, the critical determinant of issay sensitivity; this parameter is essentially overned by the precision of the density measure-

From Eq. (1), the fraction of sites unoccupied y analyte = 1/(1 + p), and the fraction occupied y anti-idiotypic antibody = Q/(1 + p)(1 + Q). hus, if the CV in the measurement of antiliotypic antibody is ε , the standard deviation is 2/(1+p)(1+Q). This term also represents the D in the estimate of the fraction of sites occupied y analyte. Since the total number of antibody nding sites in the spot is DA, the SD in the timate of occupied sites as $p \to 0$ (i.e. σD_0^*) proximates $\varepsilon DAQ/(1 + Q)$; the SD in the cupied site surface-density estimate is thus Q/(1+Q). But the SD in the measurement of actional binding-site occupancy when $p \rightarrow 0$ fines D_{\min} , and hence the minimum detectable alyte concentration in the test sample as dicated in Eq (4). Thus

$$D_{\min} = D_{\min} \times [(6 \times 10^{20})(1 + Q)]/DKQ$$
 (5)

$$= \varepsilon DQ/(1+Q) \pm [(6 \times 10^{20})(1+Q)] DKQ$$
 (6)

$$= \varepsilon/K \times (6 \times 10^{20}) \tag{7}$$

For example, if values of Q = 1, $D = 10^5$ molecules/ μ m², and $K = 10^{11} L/M$ are assumed as in the non-competitive example considered above, and the CV in the measurement of anti-idiotypic antibody density in the microspot is 1% (i.e. $\varepsilon = 0.01$), then $D_{\min} = 500$ molecules/ μm^2 , and $C_{\min} = 6 \times 10^7$ molecules/ml = 10-13 M/L. Fractional occupancy of the sensing antibody binding sites by the minimum detectable analyte concentration is, in this example, 1%. It should be noted that the sensitivity limit of ϵ/K (expressed in molar terms) is identical to that previously established for conventional competitive' assays (Ekins and Newman, 1970), and which underlies the predictions represented in Fig. 4.

Such considerations appear to suggest (a) that microspot assay sensitivities superior to those obtainable by conventional radioisotopically based immunoassays are achievable, and (b) that sensitivities yielded by non-competitive microspot assays are likely to be considerably greater than those of corresponding competitive microspot assays. It must be emphasized, however, that, though such predictions are likely to prove correct, assumptions regarding the performance of the labels and signal-measuring instrument used are incorporated in the simple theoretical analysis discussed above. Such factors are clearly of importance in determining overall microspot immunoassay performance.

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Practical implementation

The concepts discussed above are clearly exploitable using a variety of antibody labels, including chemiluminescent labels; however, our preliminary studies have been based on the use of conventional fluorophores, since the technology of simultaneous measurement of dual fluorescence from small areas is already well established. Because this volume centres on chemiluminescence, we shall provide only a brief indication of our initial experimental work in this area, which is currently based on the use of commercially available confocal microscopes.

Instrumentation: the laser scanning confocal microscope. In laser scanning confocal fluoresc-

Additional States ence microscopy, a small area of the specimen is illuminated by a focused laser beam; the fluorescence photons emanating solely from this area are, in turn, focused onto a photon detector. Both the intensity of illumination and the efficiency of light collection diminish rapidly with distance from the focal plane (Fig. 12). At the 'confocal' point, the projection of the illumination pinhole and the back-projection of the detector pinhole coincide. Such systems contrast with conventional epifluorescence methods, where the specimen is exposed to an essentially uniform flux of illumination (White et al., 1987).

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Exercise 4

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Sensitivity of current instruments. Typically, fluorescence photons emanating from the laser-

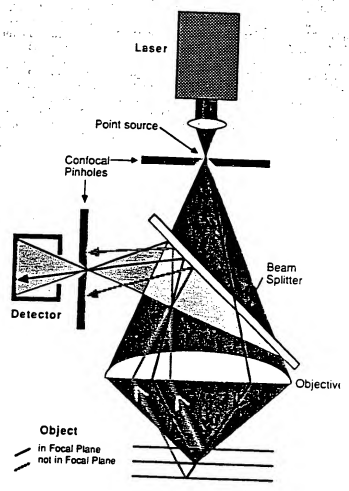


Figure 12. Principle of the confocal microscope. Illuminating light is focused at a point in the focal plane. Reflected light from this point is focused onto a detector. A complete two-dimensional image of structures within the focal plane is obtained by scanning the selected area of interest, and may be stored in a microcomputer for video display

illuminated area are detected by a low darkcurrent photomultiplier. Electrons spontaneously emitted by the photomultiplier photocathode contribute to the background signal of the instrument, and must, for highest sensitivity, be minimized. Fortunately the overall design of such instruments permits the photomultiplier photocathode to be of very small area; so that this particular source of background noise is not only small, but can be expected to reduce in relative importance with future improvement in photomultiplier design. Meanwhile current instruments already display very high sensitivity of detection of fluorescent signals. For example, the confocal microscope manufactured by Zeiss is claimed to display a lower detection limit for fluorescein of about ten molecules/µm² (Ploem, 1986). Most commercially available FITC-labelled IgG attains a fluorophore/protein molar ratio of ~4; thus the detection limit (D_{\min}^*) of the Zeiss microscope is ~2-3 FITC-labelled IgG molecules/µm². This implies an analyte-concentration detection limit of $\sim 2.4 \times 10^5$ molecules/ml for a two-site assay, assuming the same parameter values as used in the examples discussed above, or 2.4×10^4 molecules/ml using a 'sensing' antibody of affinity $10^{12} L/M$.

Another comparable instrument is the Bio-Rad/Lasersharp laser scanning confocal microscope, which we are currently using in the development of 'ratiometric' multi-analyte assay methodology in accordance with the principles outlined above (see Fig. 13). The argon laser in this system possesses two excitation lines at 488 and 514 nm. It is thus particularly efficient for the excitation of blue/green emitting fluorophores such as FITC (which displays an excitation maximum at 492 nm). However, it is considerably less efficient in the excitation of red-emitting fluorophores such as Texas red (excitation maximum 596 nm). However, the ratiometric immunoassay principle permits considerable variation in detection efficiencies of the two labels relied on since, inter alia, the specific activities of the two labelled antibody species forming the antibody couplets can be chosen to yield optimal signal ratios in the region of unity. Thus inefficiency of the argon laser in exciting red emitting fluorophores is not necessarily a major handicap in the present context.

Though the current Lasersharp instrument relies on a conventional microscope rather than a purpose-designed optical system (and appears to

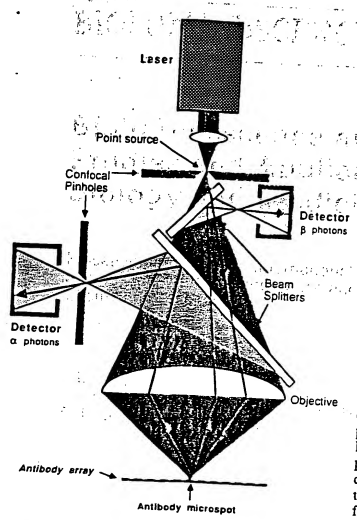


Figure 13. Dual-channel confocal fluorescence microscope permitting simultaneous measurement of the fluorescence signals from two fluorophors situated at the focal point. By scanning the antibody array, the ratio of signals from each antibody microspot may be determined

be less sensitive), it permits quantification of fluorescence signals generated from microspots of selected area. Initial studies have revealed that, under conditions that are not necessarily optimal, the instrument is capable of detecting approximately twenty-five FITC-labelled IgG molecules/

µm², scanning an area of ~50 µm² (Fig. 14). It must be stressed that neither of these confocal microscopes are designed specifically for routine ratiometric multi-analyte immunoassay use, and it can be anticipated that future instruments constructed specifically for this purpose are likely to prove both cheaper and more sensitive.

Other instruments. The MPM 200 Microscope Photometer manufactured by Zeiss of West

Germany is anticipated to become available shortly. This photometer is claimed to be highly versatile: it can be used in transmission and reflection modes, and as a highly sensitive fluorimeter. The measuring field can be varied in shape and size for optimum adjustment to the specimen structure. More generally, the technology of sensitive light measurement is improving rapidly in response to needs in astronomy, the space program etc., such technology clearly being readily exploitable in a multi-analyte immunoassay context using light-generating labels in accordance with the broad principles presented here.

Solid antibody supports. On the basis of the theoretical considerations discussed above, it is evident that solid antibody supports for multianalyte immunoassay use should display a capacity to adsorb a high surface density of antibody combined with low intrinsic signal-generating properties (for example, low intrinsic fluorescence), thus minimizing background. We have examined a number of materials, including polypropylene, Teflon, cellulose and nitrocellulose membranes and microtitre plates (clear polystyrene plates from Nunc; black, white and clear polystyrene plates from Dynatech withthese criteria in mind. White Dynatech Microfluor microtitre plates, formulated specially for the detection of low fluorescence signals, yield high signal-to-noise ratios and have therefore been provisionally used in our developmental studies.

Surface density of antibody coating. Preliminary experiments using Microfluor plates have revealed that it is possible to coat them with antibody at a surface density of at least 5×10^4 IgG molecules/ μ m² (Fig. 15). Moreover nearly all antibody molecules so deposited appear to retain immunological activity (Fig. 16).

Verification of the 'ratiometric' imunoassay concept. Our primary intention, in initial studies, has been establishment of the basic conditions which, using a particular instrument, can be anticipated on theoretical grounds to yield high assay sensitivity. Though the setting up of individual microspot immunoassays has thus appeared to us to be of secondary importance during the initial stages of our studies, we have nevertheless

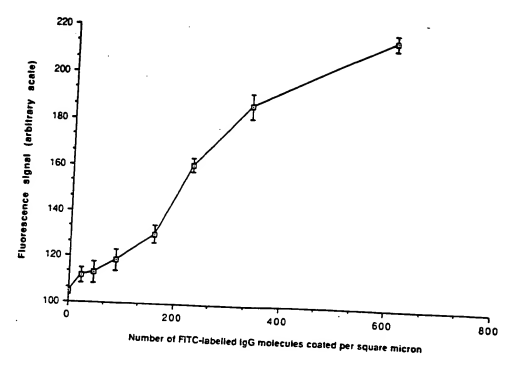


Figure 14. Fluorescence signal (arbitrary units), measured using the Bio-Rad/Lasersharp scanning confocal microscope, plotted as a function of the density of fluorescein-labelled IgG molecules (number of molecules/μm²) depositied on Dynatech Microfluor white microtitre plates

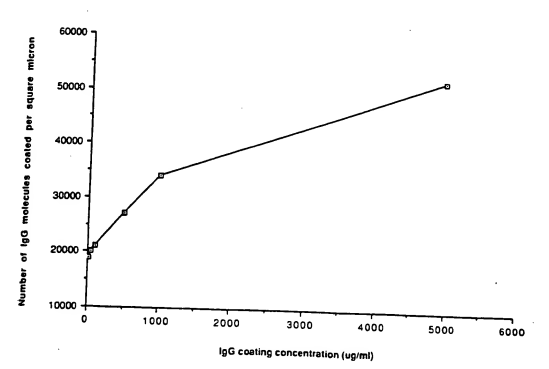


Figure 15. Surface density of IgG molecules (number of molecules/ μ m²) deposited on Dynatech Microfluor white plates plotted as a function of IgG concentration (μ g/ml) in the coating solution

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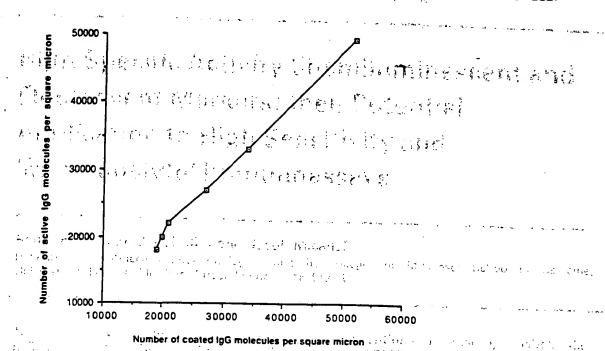


Figure 16. Surface density of immunoreactive IgG molecules (number of molecules/ μ m²) plotted as a function of the total surface density of IgG (number of molecules/ μ m²) on Dynatech Microfluor white microtitre plates

thought it useful to confirm the validity of our general concepts by comparing the performance of certain assays when constructed in microspot format and when conventionally designed. For example, we have compared a dual-labelled tumour necrosis factor (TNF) ratiometric assay system using Texas red and FITC-labelled antibodies with an optimized IRMA system using identical antibodies but with the second antibody 1251-labelled. Although unoptimized, the ratiometric microspot assay yielded formal sensitivity values closely approaching that of the conventional, optimized, IRMA. Although verifying the general concepts underlying ratiometric microspot immunoassay methodology, further work is required to achieve the considerably greater sensitivity that theory predicts as achievable using optimized reagent concentrations and improved instrumentation.

CONCLUSION

As indicated above, differentiation of the fluorescent signals yielded by two fluorophores can be readily achieved solely on the basis of wavelength differences, and this approach has been relied on entirely in our preliminary studies. However,

other physical techniques exploiting differences in decay time of two or more fluorescence emissions (using, for example, a pulsed or sinusoidally modulated laser source, and time- or phaseresolving detectors) are available, and can be expected both to further reduce background and to improve signal resolution, thus increasing assay sensitivity and precision. These considerations aside, the basic technology involved closely resembles that employed in domestic compact. disk recorders and other similar data-storage devices, the obvious difference being that light emitted from each of the discrete zones forming the antibody-array is fluorescent rather than reflected, and yields chemical rather than physical information. Indeed, our preliminary studies suggest that highly sensitive immunoassays using antibody microspots of surface area approximating 50 µm² are achievable, implying that some 2,000,000 different immunoassays could, in principle, be accommodated on a surface area of 1 cm². Though non-specific binding of a multiplicity of developing antibodies would probably prohibit the use of antibody arrays of this order, it is evident that the technology is capable of encompassing analyte numbers of the kind likely to be useful in practice.

The development of multi-analyte assay systems of this kind can be anticipated to bring about

fundamental changes in medical diagnosis and Ekins, R. P. (1980). More sensitive immunoassays. Nature, many other biologically related areas. Systems capable of measuring every hormone and other endocrinologically related substance within a single small sample of blood are within technological reach, providing data which, when analysed with the aid of computer-based 'expert' patternrecognition systems, are likely to reveal endocrine deficiences only dimly perceived using current 'single-analyte' diagnostic procedures. Such systems also provide a means to the development of a 'random access' immunoassay methodology, permitting the selection of any particularized test, or combination of tests from an extensive analyte menu. Clearly the accommodation of a wide range of individual immunoassays on a small immunoprobe (comparable in its overall physical dimensions with a few drops of blood) is likely to totally transform the logistics of his immunodiagnostic testing, and genuinely represents, in our view, 'next generation' immunoassay methodology.

Acknowledgement

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REFERENCES

- Barnard, G. J. R., Kim, J. B. and Williams, J. L. (1985). Chemiluminescence immunoassays and immunochemiluminometric assays. In Alternative Immunoassays, Collins, W. P. (Ed.), John Wiley, Chichester, pp. 123-152.
- Berson, S. A. and Yalow, R. S. (1973). Measurement of hormones-radioimmunoassay. In Methods in Investigative and Diagnostic Endocrinology, 2A, Berson, S. A. and Yalow, R. S. (Eds), North Nolland/Esevier, New York,
- Dakubu, S., Ekins, R., Jackson, T. and Marshall, N. J. (1984). High sensitivity, pulsed light time-resolved fluoroimmunoassay. In Practical Immunoassay. The State of the Art, Butt, W. R. (Ed.), Marcel Dekker, New York, pp. 71-101.
- Ekins, R. P. (1976). General principles of hormone assay. In Hormone Assays and their Clinical Application, Loraine, J. A. and Bell, E. T. (Eds), Churchill Livingstone, Edinburgh, pp. 1-72.
- Ekins, R. P. (1978). The future development of immunoassay. In Radioimmunoassay and Related Procedures in Medicine 1977, IAEA, Vienna, pp.241-275.
- Ekins, R. P. (1979). Radioassay methods. In Radiopharmaceuticals II: Proceedings, 2nd International Symposium on Radiopharmaceuticals, 19-22 March 1979, Seattle, Washington, Sorenson, J. A. (Ed), Society of Nuclear Medicine, New York, pp. 219-240.

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- Ekins, R. P. (1983a). The precision profile: its use in assay design, assessment and quality control. In Immunoassays for Clinical Chemistry, Hunter, W. M. and Corrie, J. E. T. (Eds), Churchill Livingstone, Edinburgh, pp.:76-105.
- Ekins, R. P. (1983b). Measurement of analyte concentration. British Patent no. 8224600.
- Ekins, R. (1985). Current concepts and future developments. In Alternative Immunoassays, Collins, W. P. (Ed.), John Wiley, Chichester, pp. 219-237.
- Ekins, R. P. and Newman, B. (1970). Theoretical aspects of saturation analysis. In Karolinska Symposia on Research Methods in Reproductive Endocrinology. 2nd Symposium: Steroid Assay by Protein Binding, Diczfalusy, E. (Ed.), The Reproductive Endocrinology Research Unit, Karolinska sjukhuset Stockholm, pp. 11-36.
 - Ekins, R. P., Newman, B. and O'Riordan, J., L., H. (1968). Theoretical aspects of 'saturation' and radioimmunoassay. In Radioisotopes in Medicine: In Vitro Studies, Hayes, R. L., Goswitz, F. A. and Murphy, B. E. P. (Eds), Oak Ridge Symposia, USAEC, Oak Ridge, Tennessee, pp.
- Ekins, R. P., Newman, B. and O'Riordan, J. L. H. (1970a). Saturation assays. In Statistics in Endocrinology, Me-Arthur, J. W. and Colton, T. (Eds), MIT Press, Cambridge, MA, pp. 345-378.
- Ekins, R. P., Newman, B. and O'Riordan, J. L. H. (1970b). Competitive protein-binding assays. Discussion. In Statistics in Endocrinology, McArthur, J. W. and Colton, T. (Eds), MIT Press, Cambridge, MA, pp. 379-392.
- Ekins, R. P., Filetti, S., Kurtz, A. B. and Dwyer, K. (1980). A simple general method for the assay of free hormones (and drugs); its application to the measurement of serum free thyroxine levels and the bearing of assay results on the 'free thyroxine' concept. J. Endocrinol., 85, 29-30.
- Ezzell, C. (1986). Hybritech versus Abbott. Nature, 324, 506. Ezzell, C. (1987a). Judge confirms injunction in sandwich assay patent suit. Nature, 326, 532.
- Ezzell, C. (1987b). Hybritech wins court injunction over sandwich assays. Nature, 327, 5.
- Hemmilä, I., Dakubu, S., Mukkala, V.-M., Siiteri, H. and Lovgren, T. (1983). Europium as a label in time-resolved immunosluorometric assays. Anal. Biochem., 137, 335-
- Jackson, T. M., Marshall, N. J. and Ekins, R. P. (1983). Optimisation of immunoradiometric (labelled antibody) assays. In Immunoasays for Clinical Chemistry, Hunter, W. M. and Corrie, J. E. T. (Eds), Churchill Livingstone, Edinburgh, pp. 557-575.
- Kohen, F., Bayer, E. A., Wilchek, M., Barnard, G., Kim, J. B., Collins, W. P., Beheshti, I., Richardson, A. and McCapra, F. (1984). Development of luminescence-based immunoassays for haptens and for peptide hormones. In Analytical Applications of Bioluminescence and Chemiluminescence, Kricka, L., Stanley, P. E., Thorpe, G. H. G. and Whitehead, T. P. (Eds), Academic Press, New York, pp. 149-158.
- Kohen, F., Pazzagli, M., Serio, M., DeBoever, J. and Vanderkerckhove, D. (1985). Chemiluminescence and bioluminescence immunoassay. In Alternative Immunoassays, Collins, W. P. (Ed). John Wiley, Chichester, pp. 103-121.
- Köhler, G. and Milstein, C. (1975). Continuous culture of

- fused cells secreting specific antibody. Nature, 256, 495-497.
- Loevinger, R. and Berman, M. (1951). Efficiency criteria in radioactive counting. *Nucleonics*, 9, 26:
- Marshall, N. J., Dakubu, S., Jackson, T. and Ekins, R. P. (1981). Pulsed-light, time-resolved, fluoroimmunoassay. In Monoclonal Antibodies and Developments in Immunoassay, Albertini, A. and Ekins, R. (Eds), Elsevier/North Holland, Amsterdam, pp. 101-108.
- McCapra, F., Tutt, D. E. and Topping, R. M. (1977). Assay method utilizing chemiluminescence. British Paient no. 1, 461–877
- McGown, L. B. and Bright, F. V. (1984). Phase-resolved in fluorescence spectroscopy. Anal. Chem., 56, 1400-1417.
- Miles, L. E. H. and Hales, C. N. (1968). An immunoradiometric assay of insulin. In Protein and Polypeptide Hormones, Pt. 1, Margoulies, M. (Ed.), Excerpta Medica, Amsterdam, pp. 61-70.
- Ploem, J. S. (1986). New instrumentation for sensitive image analysis of fluorescence in cells and tissues. In Applications of Fluorescence in the Biological Sciences, Tayer, D. L., Waggoner, A. S., Lanni, F., Murphy, R. and Birge, R. (Eds), Alan R. Liss, New York, pp. 289-300.
- Rodbard, D. and Weiss, G. H. (1973). Mathematical theory of immunometric (labelled antibody) assay. *Analys. Biochem.*, 52, 10-44.
- Shalev, A., Greenberg, G. H. and McAlpine, P. J. (1980). Detection of attograms of antigen by a high sensitivity enzyme-linked immunosorbent assay (HS-ELISA) using a fluorogenic substrate. J. Immunol. Methods, 38, 125-139.
- Tait, J. F. (1970). Competitive protein-binding assays. Discussion. In Statistics in Endocrinology, McArthur, J. W. and Colton, T. (Eds), MIT Press, Cambridge, MA. pp. 379-392.

- Weeks, I., McCapra, F., Campbell, A. K. and Woodhead, J. S. (1983). Immunoassays using chemiluminescent labelled antibodies. In *Immunoassays for Clinical Chemistry*, Hunter, W. M. and Corrie, J. E. T. (Eds.), Churchill
- Livingstone, Edinburgh, pp. 525-530.

 Weeks I. Campbell A K. Woodbead S. and McConn. F.
- Weeks, I., Campbell, A. K., Woodhead, S. and McCapra, F. (1984). Immunoassays using chemiluminescent labels. In Practical Immunoassay. The State of the Art, Butt, W. R. (Ed.), Marcel Dekker, New York, pp. 103-116.
- White, J. G., Amos, W. B. and Fordham, M. (1987). An evaluation of confocal versus, conventional imaging of biological structures by fluorescence light microscopy. J. Cell Biol., 105, 41-48.
- Whitehead, T. P., Thorpe, G. H., Carter, T. J., Groucutt, C. and Kricka, L. J. (1983). Enhanced luminescence procedure for sensitive determination of peroxidase-labelled conjugates in immunoassay. *Nature*, 305, 158-159.
- Wide, L. (1971). Solid phase antigen-antibody systems. In Radioimmunoassay Methods. Kirkham, K. E. and Hunter, W. M. (Eds), Churchill Livingstone, Edinburgh, pp. 405-418.
- Woodhead, J. S., Addison, G. M., Hales, C. N. and O'Riordan, J. L. H. (1971). Discussion. In Radioimmuno-assay Methods, Kirkham, K. E. and Hunter, W. M. (Eds), Churchill Livingstone; Edinburgh, pp. 467-488.
- Yalow, R. S. and Berson, S. A. (1970a). Radioimmunoassays. In Statistics in Endocrinology, McArthur, J. W. and Colton, T. (Eds), MIT Press, Cambridge, MA. pp. 327-344.
- Yalow, R. S. and Berson, S. A. (1970b). Competitive protein-binding assays. Discussion. In Statistics in Endocrinology. McArthur, J. W. and Colton, T. (Eds.), MIT Press, Cambridge, MA. pp. 379-392.

JOURNAL OF _____BIOLUMINESCENCE AND CHEMILUMINESCENCE

Bioluminescence and Chemiluminescence: Studies and Applications in Biology and Medicine

Proceedings of the Vth International Symposium on Bioluminescence and Chemiluminescence

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Editors:

M. Pazzagli, E. Cadenas, L. J. Kricka, A. Roda and P. E. Stanley

Volume 4 1989



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CLN. CHEM. 37/11, 1955-1967 (1991) rain the second of the second Multianalyte Microspot Immunoassay Microanalytical "Compact Disk" of the Future R. P. Ekins and F. W. Chu per stially to produce the product of the product o

16. [16] 、 在有些在一点,不是在一个情况。 Throughout the 1970s, controversy centered both on immunoassay "sensitivity" per se and on the relative sensitivities of labeled antibody (Ab) and labeled analyte methods. Our theoretical studies revealed that RIA sensitivities could be surpassed only by the use of very high-specificactivity nonisotopic labels in "noncompetitive" designs, preferably with monoclonal antibodies. The time-resolved fluorescence methodology known as DELFIA—developed in collaboration with LKB/Wallac-represented the first commercial "ultrasensitive" nonisotopic technique based on these theoretical insights, the same concepts being subsequently adopted in comparable methodologies relying on the use of chemiluminescent and enzyme labels. However, high-specific-activity labels also permit the development of "multianalyte" immunoassay systems combining ultrasensitivity with the simultaneous measurement of tens, hundreds, or thousands of analytes in a small biological sample. This possibility relies on simple, albeit hithertounexploited, physicochemical concepts. The first is that all immunoassays rely on the measurement of Ab occupancy by analyte. The second is that, provided the Ab concentration used is "vanishingly small," fractional Ab occupancy is independent of both Ab concentration and sample volume. This leads to the notion of "ratiometric" immunoassay, involving measurement of the ratio of signals (e.g., fluorescent signals) emitted by two labeled Abs, the first (a "sensor" Ab) deposited as a microspot on a solid support, the second (a "developing" Ab) directed against either occupied or unoccupied binding sites of the sensor Ab. Our preliminary studies of this approach have relied on a dual-channel scanning-laser confocal microscope, permitting microspots of area 100 μm^2 or less to be analyzed, and implying that an array of 10⁶ Ab-containing microspots, each directed against a different analyte, could, in principle, be accommodated on an area of 1 cm2. Although measurement of such analyte numbers is unlikely ever to be required, the ability to analyze biological fluids for a wide spectrum of analytes is likely to transform immunodiagnostics in the next decade.

Additional Keyphrases: ratiometric Immunoassays . scanninglaser confocal microscope · fluoroimmunoassay

Immunoassay and other protein-binding assay methods based on the use of radioisotopic labels have played a major role in medicine during the past three decades.

Department of Molecular Endocrinology, University College and Middlesez School of Medicine, Mortimer St., London WIN

Presented at the 23rd annual Oak Ridge Conference on Advanced Analytical Concepts for the Clinical Laboratory, St. Louis, MO, April 1991.

Take pages 100 Take believe the first his page. Their utility and importance have derived primarily from the structural specificity of many reactions between binding proteins and analytes and the detectabile ity of isotopically labeled reagents, the latter endowing such techniques with "exquisite sensitivity." Recently, however, interest has increasingly focused on nonisotopic techniques based on identical analytical principles, differing only in the nature of the marker used to label the reactant (e.g., antibody or antigen), whose distribution between reacted ("bound") and unreacted ("free") fractions constitutes the assay "response."

The basic aims underlying this interest can be broadly classed under four main headings:

· avoidance of the environmental, legal, economic, and practical disadvantages of isotopic techniques (e.g., limited shelf life of isotopically labeled reagents, problems of radioactive waste disposal, cost and complexity of radioisotope counting equipment), particularly those impeding the development of, for example, simple diagnostic kits for home or doctor's office use;

• achievement of greater assay sensitivity; "direct" measurement of analyte concentrations by use of transducer-based "immunosensors";

· simultaneous measurement of multiple analytes ("multianalyte assay").

In this presentation I will focus primarily on the last of these objectives, using this to set out the principles underlying our present attempts to develop a new "ministurized" technology that will permit the simultaneous measurement of an unlimited number of analytes in a small biological sample such as a single drop of blood. However, retention (and, if possible, improvement) of the high sensitivities of conventional isotopic techniques is a basic aim not only of our own studies in this area but also of most other endeavors falling under the above headings. It is therefore appropriate to preface this paper with a discussion of the general principles underlying the attainment of high binding-assay sensitivity.

immunoassay Sensitivity: Some Basic Concepts Definition of Assay Sensitivity

The need to establish assay conditions yielding maximal sensitivity underlay the independent construction of mathematical theories of immunoassay design by both Yalow and Berson (1) and Ekins et al. (2) in the course of the original development of these methods in the early 1960s. Regrettably, these theoretical studies led to a prolonged controversy, arising largely from the conflicting concepts of "sensitivity" adopted by the two groups (see Figure 1). Briefly, Berson and Yalow, in their many publications relating to immunoassay design (e.g., 1, 3), defined sensitivity as the slope of the

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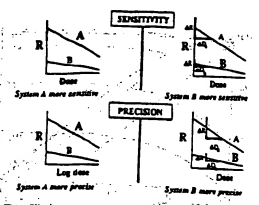


Fig. 1. The differing concepts of sensitivity and precision underlying radioimmunoassay design theories developed by (left) Yalow and Berson (e.g., 1, 3) and (right) Ekins et al. (2, 4) Yalow and Berson define assay A as more sensitive because it yields a response curve of greater stope. Ekins et al. define assay B as more sensitive because the imprecision of measurement of zero dose $(\sigma_{\rm D})$ is less. Yalow and Berson likewise define an assay system as more precise if it yields a steeper response curve when data are plotted on a tog dose scale

response curve relating the fraction or percentage of labeled antigen bound (b) to analyte concentration ([H]). In contrast, Ekins et al. (e.g., 2, 4) defined sensitivity as the (im)precision of measurement of zero dose, this quantity being indicative of, and essentially equivalent to, the lower limit of detection.

The key difference between these two definitions clearly lies in the dependence of the assay detection limit on the error (imprecision) in the measurement of the response variable. By neglecting this crucial factor, the "response curve slope" definition leads to many obvious absurdities. For example, plotting conventional RIA data in terms of the response metameter B/F (i.e., the bound to free ratio) suggests that assay "sensitivity" is increased by increasing the antibody concentration in the system; however, the converse conclusion is reached if identical data are plotted in terms of F/B (see Figure 2). Observation of the shape and slopes of response curves without detailed error analysis thus constitutes a totally misleading guide to optimal immunoassay design. This approach has, however, characterized many of the studies conducted in the immunoassay field during the past 30 years, and has been the source of much

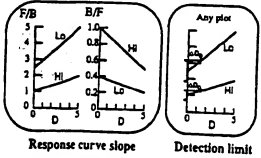


Fig. 2. Schematic representation of RIA dose-response curves observed for high and low antibody concentrations plotted in terms of (left) the free/bound fraction (F/B); (center) the bound/free fraction (B/F)

Note that the low antibody concentration yields a response curve of greater slope when the assay response is plotted in terms of F/B, but of lower slope when plotted in terms of B/F. The precision of measurement of zero dose (ΔD_0) is independent of the coordinate frame used to plot assay data (see right)

mythology. For example, consideration of the Law of Mass Action reveals that, when response curves corresponding to different antibody concentrations are plotted in terms of b vs [H], the maximal alope at zero dose is obtained for a concentration of 0.5/K (where K is the affinity constant), in which circumstance the zero dose response (b₀) is 33%. This conclusion led to Berson and Yalow's enunciation of the well-known dictum (which, albeit erroneous, is broadly adhered to by many immunoassay practitioners and kit manufacturers) that, to maximize RIA sensitivity, the amount of antibody to use in the system is that which binds 33% of labeled antigen in the absence of unlabeled antigen (1, 3).

Disagreement regarding the concept of sensitivity inevitably led to prolonged dispute regarding immunoassay design (5). However, although it is still common to encounter publications in the field that rely solely on the response curve slope as a measure of sensitivity, the assay detection limit is now widely accepted as the only valid indicator of this parameter, and we do not therefore intend to dwell further on this issue here. It is nevertheless relevant to an understanding of the "miniaturized" assay methodology described below to emphasize that untenable concepts of both sensitivity and precision underlie many of the commonly accepted rules governing current immunoassay-design practice, some of which are contravened in our own approach.

Basic Immunoassay Designs

It is likewise important in the present context to comprehend the basis of the various types of immunoassays currently in use, and the constraints on the sensitivities of which they are potentially capable. The radio-immunoassay and analogous protein-binding assay techniques originally developed for the measurement of insulin by Yalow and Berson (6), and of thyroxin and vitamin B₁₂ by Ekins and Barakat (7, 8), relied on the use of a labeled analyte marker to reveal the products of the binding reactions between analyte and binder (Figure 3, left). This approach has subsequently often been portrayed as relying on "competition" between labeled and unlabeled analyte molecules for a limited number of protein-binding sites, such assays being frequently referred to as "competitive."

Subsequently, Wide et al. in Sweden (9), followed shortly by Miles and Hales in the U.K. (10), developed labeled antibody methods (Figure 3, right). These methods represented an extension of the "labeled reagent" methods (utilizing radiolabeled organic compounds such as ¹³¹I-labeled p-iodosulfonyl chloride, [³H]acetic anhydride, and other similar reagents) devised, during the early 1950s, by Keston et al. (11), Avivi et al. (12), and others for quantifying amino acids, steroid and thyroid hormones, etc. Although radiolabeled antibody methods (immunoradiometric assays; IRMAs) were originally claimed (13) to be more sensitive than methods based on the use of radiolabeled analyte, these claims were supported by neither rigorous theoretical analysis nor persuasive experimental evidence, and for some time remained controversial. Further doubt on their validity

Fig. 3. Labeled-analyte (left) and labeled-antibody (right) assay systems compared

Labeled-analyte assay systems assentially rely on observation of an analyte "marker" to reveal the products of the reaction between analyte and antibody (although the labeled analyte is not necessarily identical to the unlabeled analyte in its blinding characteristics vis-d-vis antibody). Note that, irrespective of which fraction of the labeled analyte is measured after the binding reaction, the optimal antibody concentration required to maximize sensitivity in such a system tends toward zero (assuming a background signal of 0). Labeled-antibody systems rely on observation of an antibody "marker" to reveal the products of the binding reaction between analyte and antibody. In this case, the optimal antibody concentration required to maximize sensitivity tends toward zero when the "tree" antibody fraction is measured, but tends toward infinity when the bound fraction is determined (likewise assuming zero background)

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was cast by the publication by Rodbard and Weiss in 1973 (14) of detailed theoretical studies demonstrating that both labeled analyte and labeled antibody methods possessed essentially equal sensitivities. (Note: These authors suggested that IRMAs might be more sensitive in the assay of small polypeptides, in which radioiodine incorporation into the antigen molecule was restricted; conversely, these assays would be less sensitive for the measurement of antigens of high molecular mass.) Nevertheless, despite the appearance of this publication, the belief that labeled antibody methods per se are intrinsically more sensitive than the corresponding labeled analyte methods gained wide acceptance among clinical chemists.

The reason for confusion on this issue is that the greater potential sensitivity of certain assay formats is not really a consequence of the labeling of antibody as opposed to analyte; indeed, the apparent antithesis between labeled-analyte and labeled-antibody methods diverts attention from the true reasons underlying the superior sensitivity of certain assay designs. Theoretical analysis (see, e.g., 4, 15) reveals that, assuming "perfect" separation of the products of the binding reaction (i.e., no misclassification of bound and free moieties), the optimal antibody concentration (for maximal sensitivity) in a labeled analyte immunoassay invariably tends to zero, irrespective of whether the free or bound labeled analyte fraction is measured, whereas in labeled-antibody methods the optimal antibody concentration depends on which labeled-antibody fraction is measured (see Figure 3). If the free (unreacted) antibody fraction is measured, the optimal concentration also tends to zero; conversely, if the analyte-bound fraction is measured, the concentration tends to infinity. In short, of the four basic measurement strategies available—labeled analyte, with measurement of free or bound reaction product, and labeled antibody, also with measurement of free or bound product—only one permits, in practice, the use of antibody concentrations approaching infinity.

This particular approach may, for want of a better term, be described as "noncompetitive," although it must be emphasized that such terminology involves a departure from the original meanings attached to "competitive" and "noncompetitive" when these descriptions were first used in the present context. Indeed, as discussed below, assays may be subclassified in this manner when no labeled reagent of any kind is involved.

However, the categorization of immunoassays and other binding assays as competitive or noncompetitive, depending on the binding agent concentration yielding maximal assay sensitivity, itself obscures the underlying reasons for the existence of this divergence in assay designs, and may thus be misleading. These reasons may be more readily understood if the basic principles of such assays are portrayed differently from their customary presentation.

The "Antibody Occupancy Principle" of Immunoassay

When a "sensor" antibody is introduced into an analyte-containing medium, binding sites on the antibody are occupied by analyte molecules to a fractional extent that reflects both the equilibrium constant governing the binding reaction, and the final concentration of free analyte present in the mixture. This proposition stems immediately from the Law of Mass Action, which can be written as

$$[AbAg][[Ab] = K[fAg]$$

or as fractional occupancy of antibody binding sites,

$$[AbAg]/[Ab] = K[fAg]/(1 + K[fAg])$$
 (2)

where [AbAg], [Ab], [fAb], and [fAg] represent the concentrations (at equilibrium) of bound and total antibody, and free antibody and antigan (analyte), respectively, and K = equilibrium constant. The final concentration of free analyte generally depends on the concentrations of both total analyte and antibody; however, when total antibody approximates 0.05/K or less, free and total antigen ([Ag]) concentrations do not differ significantly, and fractional occupancy of antibody is given by

$$[AbAg]/[Ab] = K[Ag]/(1 + K[Ag])$$
 (3)

Assays utilizing this concept have been termed "ambient analyte immunoassays" (16), fractional occupancy being independent of both sample volume and antibody concentration (see below).

All immunoassays essentially depend on measurement of the "fractional occupancy" of the sensor antibody after its reaction with analyte (see Figure 4). Techniques relying on the measurement of unoccupied antibody binding sites (from which antibody occupancy is implicitly deduced by subtraction) necessitate—for attainment of maximal sensitivity—the use of sensor antibody concentrations tending to zero; these assays

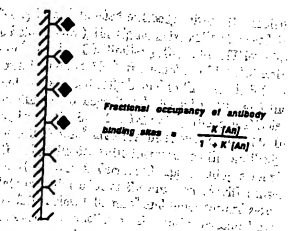


Fig. 4. The antibody binding-site occupancy principle of immunoas-

All immunoassays implicitly rely on the measurement of (tractional) bindingsite occupancy by analyte

may therefore be categorized as "competitive." Conversely, techniques in which occupied sites are directly measured permit (in principle) the use of relatively high concentrations of sensor antibody and may be described as "noncompetitive." This difference in assay design simply reflects the proposition that, to minimize error in the measurement, it is generally undesirable to measure a small quantity by estimating the difference between two large quantities.

These concepts are illustrated in Figure 5, which portrays basic immunoassay formats currently in common use. Conventional RIA and other similar "labeledanalyte" techniques rely on measurement of unoccupied binding sites, generally by back-titration (either simultaneous or sequential) with labeled analyte, but antiidiotypic antibody (reactive only with unoccupied sites on the sensor antibody) may be used for the same purpose. In the case of single-site labeled-antibody assays, the labeled antibody itself constitutes the sensor antibody; after reaction with analyte, this sensor antibody may be separated into occupied and unoccupied fractions through use of (e.g.) an immunosorbant (comprising antigen, antigen analog, or anti-idiotypic antibody linked to a solid support). If, after separation, the "signal" emitted by labeled antibody bound to analyte (i.e., the "occupied" fraction) is measured directly, the assay can be classed as "noncompetitive." Conversely, if one measures the labeled antibody not bound to analyte (i.e., that attached to the immunosorbant), then the assay is "competitive."

Two-site "sandwich" assays are clearly more complex because they rely on two antibodies and can be considered from two points of view. For our present purposes, the solid-phase antibody can be regarded as the "sensor" antibody, with the labeled antibody enabling the occupied sensor-antibody binding sites to be distinguished. Seen from this viewpoint, two-site assays may be classed as "noncompetitive."

These considerations emphasize that the differences in design distinguishing so-called competitive and noncompetitive methods are essentially unrelated to which

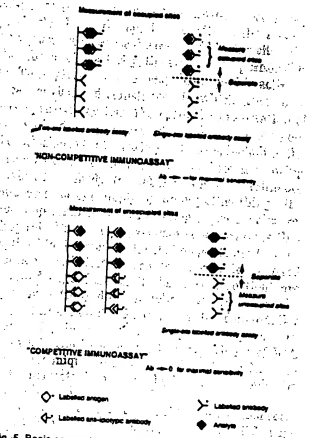


Fig. 5. Basic competitive and noncompetitive immunoassay designs. The distinction between noncompetitive and competitive immunoassays reflects the way in which antibody binding-sits occupancy is observed. Labeled-antibody methods are "noncompetitive" if occupied sites of the (labeled) antibody are directly measured, but are "competitive" (Jower right) when unoccupied sites are measured. Labeled-antigen (Jower left) or labeled-anti-idiotypic-antibody methods (Jower center) rely on measurement of eliese unoccupied by analyte, and are therefore of "competitive" design

component (if any) of the reaction system is labeled. Indeed, in the case of transducer-based "immunosensors," no component is labeled; nevertheless, the design of the immunosensor will differ significantly, depending on whether a measurable signal is yielded by occupied or unoccupied antibody binding sites situated on its surface. In short, the terms "competitive" and "noncompetitive" merely reflect alternative approaches to the determination of the occupancy of antibody binding sites and lead to differences in the optimal antibody concentration required to minimize the effects of random errors arising in the determination.

Competitive and noncompetitive immunoassays can be shown to differ significantly in many of their performance characteristics, including their sensitivities. In both types of assays, both the affinity constant (K) of the antibody and the specific activity of the label are important in determining sensitivity; however, in practice, the sensitivity of competitive assays is primarily limited by the affinity constant of the antibody, whereas the specific activity of the label is more important in noncompetitive systems. In both cases, the "experimental" or "manipulation" error in the measurement of the zero-dose response (R_0) [i.e., the relative error (σ_R/R_0) arising from pipetting and other operations, but not including the statistical signal measurement error per

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se] is of key importance in determining "potential" assay sensitivity (i.e., the sensitivity obtained by assuming the specific activity of the label to be infinite, implying zero error in signal measurement). Thus the potential sensitivity of a competitive assay can be shown to be σ_R/KR_0 , whereas that of a noncompetitive assay is given by Rook [Ab]KRo, where, in the latter case, Ro is assumed to represent the labeled antibody misclassified as bound ([bAb]_o), commonly referred to as "nonspecifically bound" antibody. Thus $R_0[Ab] = f$, the fraction of labeled antibody that is nonspecifically bound, and $R_0\sigma_{R_0}/[Ab]KR_0 = f\sigma_{R_0}/KR_0$. Assuming that the relative error (og, Ro) in the measurement of the zero-dose response is approximately identical for both competitive and noncompetitive assays, it is evident from this simple analysis that the potential sensitivity of noncompetitive methods is greater than that of competitive methods by the factor f, i.e., by the fraction of labeled antibody that is "nonspecifically bound." For example, if the nonspecifically bound fraction is 0.01%, a noncompetitive strategy is potentially capable of a sensitivity 10 000-fold greater than that of a competitive approach, other factors being equal.

These findings are summarized in Figure 6 (left), which shows the relationships between sensitivity (expressed in terms of molecules per milliliter) and anti-

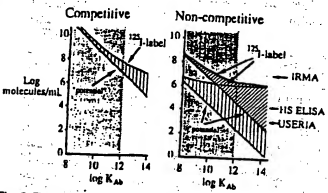


Fig. 6. Theoretically predicted sensitivities of competitive and non-competitive immunoassay methods (represented by the SD of zero analyte measurements, expressed as molecules/mL) plotted as a function of antibody affinity (K)

Note: in noncompetitive sandwich assays, the antibody affinity referred to is that of the labeled antibody. In the competitive assays, calculations are based on the assumption that the experimental error (CV) incurred in the measurement of the assay response (e.g., fraction of labeled antigen bound) is 1%. The "potential sensitivity" curve assumes the use of a label of infinite specific activity, implying that the error in the measurement of the label per se is zero. The "see class" curve indicates the loss in sensitivity arising from the statistical error incurred in counting 186 distritegrations for a finite counting time. Note that, if using antibodies with an effinity <1012 Umol (the maximum achieved in practice), little increase in sensitivity can be achieved by using labels of higher apacific activity than ¹²⁸L. For noncompetitive assays, the potential sensitivity curves shown relate to values of nonspecific binding of labeled antibody of 1% (upper curves) and 0.01% (lower curves), and emphasize the improvement in sensitivity potentially attainable by minimizing nonspecific binding. The corresponding table curves demonstrate the much greater loss in sensitivity (compared with that potentially estainable) when a radioisotopic marker is used, and the special advantages of nonisotopic labels of higher specific activity in noncompetitive assay designs (particularly if nonspecific binding is reduced to 0.1% or less). Arrows indicate assay sensitivities reported for noncompetitive immunoassays based on 120 (IRMA), and enzymes retying on fluorogenic (HS-ELISA) (28) and radioactive (USERIA) (29) substrates. These conclusions underlay the original development (19, 20) of time-resolved fluoroimmunoassay (DELFIA), the first nonisotopic "utra-aemsitive" immunoas-

body affinity in an optimized competitive (labeled analyte) assay. For this analysis, we assume (a) the use of a label of infinite specific activity, and (b) the use of 125 I as a label, the radioactivity of the samples being counted for 1 min. Computations of the theoretically optimal reagent concentrations (on which calculations represented in Figure 6 rely) were based on the further assumptions that (c) the radioactivity of the antibodybound labeled analyte fraction was counted and (d) the (relative) "experimental error" component in the measurement of the bound fraction (o,b) was 1%. Given these assumptions, the "potential" sensitivity attainable in such an assay is σ_b/K b, where K is the affinity constant of the antibody. [For example, if the affinity constant is 1012 L/mol, and o/b is 0.01 (1%), maximal assay sensitivity is 10^{-14} mol/L, or -6×10^6 molecules/ ml.] The additional "signal measurement error" arising in consequence of counting radioactive samples for a finite time implies a loss of assay sensitivity, as shown by the upper curve in Figure 6 (left). However, the resulting loss in sensitivity is relatively small for antibodies of affinities <1012 L/mol, and is negligible for antibodies with affinities <1011 L/mol. In other words, if the assayist can accept individual sample counting times of 1-5 min, little improvement in sensitivity is gained by using alternative labels of higher specific activities than 126I. However, similar considerations suggest that radioisotopic labels of much lower specific activity than 125 I (e.g., 5H) may limit the sensitivities of the assays (such as steroid assays) in which they are used, notwithstanding the use of relatively long sample counting times.

The other main conclusions stemming from such analysis are the importance of both minimizing "manipulation" errors and using antibodies of high binding affinity. For example, an increase in σ_b/b to 3% implies an approximate threefold loss in sensitivity, notwithstanding the fact that an assay reoptimized in response to the deterioration in operator skill that these numbers imply would utilize less antibody and labeled analyte, thereby partially offsetting the consequences of poor pipetting. But the most important conclusion emerging from the analysis is the near impossibility, in practice, of achieving immunoassay sensitivities better than about 107 molecules/mL by using a competitive approach, irrespective of the nature of the label used, if one assumes an upper limit to antibody binding affinities on the order of 1012 L/mol.

The results of a similar analysis of the sensitivity limitations applying to noncompetitive (two-site) assays (15) are illustrated in Figure 6 (right). Two sets of curves are portrayed here, corresponding to the assumptions of 1% and 0.01% nonspecific binding of labeled antibody to the capture-antibody substrate. Such analysis likewise yields important conclusions relevant to assay design, e.g., the crucial importance of reducing nonspecific binding of labeled antibody to an absolute minimum. Furthermore, if nonspecific binding is reduced to ~0.01%, just as high sensitivity is achievable

1 The men with the angles is the high most wife by using an antibody of $K = 10^8$ L/mol in an optimized noncompetitive assay design as by using an antibody of $K = 10^{13}$ L/mol in a competitive method. One of the most important conclusions is that the sensitivities potentially attainable with high-affinity antibodies $(K > 10^{10})$ L/mol) are beyond the reach of radioisotopically based methods, which (because of the relatively low specific activities of isotopes such as 125]) are limited in practice to sensitivities of the order of 106-107 molecules/mL or more. In short, although, under certain circumstances, noncompetitive IRMAs may be somewhat more sensitive than corresponding RIA techniques (assuming the use of the same antibody in each methodology), the potential advantages (vis-à-vis sensitivity) of the noncompetitive approach can be realized only by using nonisotopic labels of much higher specific activity than 1251. The superiority of such labels is most apparent when they are combined with high-affinity antibodies; however, Figure 6 demonstrates that, even with use of antibodies with affinities of about 10°-10° L/mol, nonisotopic labels may yield a substantial improvement in sensitivity.

These theoretical conclusions, together with the publication by Köhler and Milstein (18) of methods of in vitro production of monoclonal antibodies (1), constituted the basis of my laboratory's collaborative development (initiated around 1976) with the instrument manufacturer LKB/Wallac of the time-resolved fluorometric immunoassay methodology now known as DELFIA (19, 20). This methodology was the first "ultra-sensitive" nonisotopic immunoassay methodology to be developed. The same basic approach has subsequently been adopted by many other manufacturers, using a variety of high-specific activity labels (Table 1).

Against this background, let us now turn to the development of highly sensitive, miniaturized "microspot" immunoassays and multianalyte assay systems.

Antibody "Microspot" immunoassay: Basic Concepts

Ambient Analyte Immunoassay

Particular attention has been drawn above to the specious notion that an antibody concentration approximating 0.5/R is required to maximize the sensitivity of conventional labeled-antigen assays. This proposition is implicitly overturned by the development of "microspot" immunoassays, which we expect to provide the basis of a new generation of binding assay methods. But before

Table 1. Detection Limits According to Type of Label Specific activity

1 detectable event per second per 7.5 × 10^d labeled molecules Enzyme label Determined by enzyme "amplification factor" and detectability of reaction product Chemiluminescent label 1 detectable event per labeled molecule Fluorescent label Many detectable events per labeled molecule

Contract Contract of the discussing this methodology in detail, another basic analytical concept must be examined.

The recognition that all immunoassays essentially rely on measurement of antibody occupancy leads to a potentially important type of assay, ambient analyte immunosssay (16). This name is intended to describe assay systems that, unlike conventional methods, measure the analyte concentration in the medium to which an antibody is exposed, being independent both of sample volume and of the amount of antibody present. The possibility of developing such assays follows from the Law of Mass Action, which leads to the following equation, representing the fractional occupancy (F) by analyte of antibody binding sites (at equilibrium):

$$F^{2} - F\{(V(\underline{Ab})) + ((\underline{An} V(\underline{Ab})) + 1\} + (\underline{An} V(\underline{Ab})) = 0$$
where $(\underline{An}) = 0$ (4)

where [An] = analyte concentration, [Ab] = antibody concentration (both in units of 1/K).

From this equation it may readily be shown that, for antibody concentrations approaching 0, F = [An]/(1 +[An]). This conclusion is illustrated in Figure 7, in which the fractional occupancy of ("monospecific" or "monoclonal") antibody binding sites in the presence of various analyte concentrations is plotted against antibody concentration. When an antibody concentration of less than (say) 0.01/K (the antibody preferably, but not essentially, being coupled to a solid support) is exposed to an analyte-containing medium, the resulting (fractional) occupancy of antibody binding sites solely reflects the ambient concentration of analyte and is independent of the total amount of antibody in the system. (If, for example, $K = 10^{11}$ L/mol, an antibody binding-site concentration of 0.01/K represents $0.01 \times$ 10^{-11} mol/L, or 6.02×10^7 binding sites/mL.) Analyte binding by antibody causes depletion of (unbound) analyte in the medium but, because the amount bound is small, the resulting reduction in the ambient concentration of analyte is insignificant. For example, if the concentration of binding sites of the sensor antibodies is < 0.01/K, analyte depletion in the medium is invariably <1%, and the system is therefore effectively indepen-

The term "ambient" is used to indicate that antibody occupancy reflects the analyte concentration to which antibody binding sites are exposed, not the amount of analyte in the incubation tube; i.e., the system is independent of sample volume

Expression of reagent concentrations in terms of UK units has the affect of generalizing the graphical representation of binding assay data. The terms [Ab] and [An] are underlined to indicate that this convention has been adhered to in deriving equation 4. They do not refer to molar concentrations and are not interchangeable with [Ab] and [An]. For example, if the antibody possesses an affinity (constant) for analyte of 1011 L/mol, a concentration of 10-11 mol/L (represented in units of 1/K) is 1 (dimensionless) unit. Thus, fractional occupancy curves based on equation 4 are identical for all antibodies if this way of expressing antibody concentration is adopted: i.e., curves relating F to analyte concentration will be identical for systems using 10⁻¹¹ mol/L concentrations of an antibody with an affinity of 10¹¹ L/mol, 10⁻¹⁰ mol/L of an antibody with an affinity of 1010 L/mol, 10-9 mol/L of an antibody with an affinity of 10° L/mol, etc. (provided the analyte concentration is expressed in the same manner).

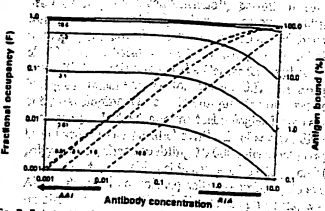


Fig. 7. Fractional antibody binding-site occupancy (F, see equation 4) plotted as a function of antibody binding-site concentration for different values of analyte (antigen) concentration (-), and the percentage binding (b) of analyte to antibody (right-hand ordinate;

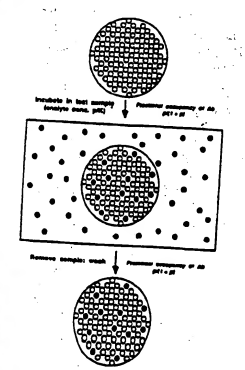
All concentrations are expressed in units of 1/K. Note that for antibody concentrations <0.01/K (approximately), the percentage binding of analyte is <1% for all analyte concentrations, and fractional binding-site occupancy is essentially unaffected by variations in antibody concentration extending over several orders of magnitude, being governed solely by antigen concentration (ambient analyte immunoassey). Note that radioimmunoasseys and other "competitive" immunoassays are conventionally designed to use antibody concentrations approximating 0.5/K-1/K or more (implying binding of analyte concentrations tending to zero (b_0) >30%), in accordance with the precepts of Yalow and Berson (1, 3)

dent of sample volume.

These conclusions lead to two further concepts. First, the antibody may be confined to a "microspot" on a solid support, such that the total number of antibody binding sites within the microspot is $< v/K \times 10^{-5} \times N$, where v the sample volume to which the microspot is exposed (in milliliters) and $N = \text{Avogadro's number } (6 \times 10^{23})$. For example, if v = 1 and $K = 10^{12}$ L/mol, then the

maximum number of binding sites that will cause negligible disturbance (<1%) to the ambient concentration of analyte is 6 x 10°, this number being greater for lower-affinity antibodies. Furthermore, the perception that the ratio of occupied (or unoccupied) sites to total binding sites is solely dependent on the ambient concentration of analyte leads to the concept of a dual-label, ratiometric," microspot immunoassay.

Dual-Label Microspot Immunoassay After exposure of a microspot of antibody (located on a suitable probe) to an analyte containing fluid (see Figure 8, left), the probe may be removed and exposed to a solution containing a high concentration of a developing" antibody directed against either a second epitope (i.e., the occupied site) on the analyte molecule if the molecule is large, or against unoccupied binding sites on the antibody in the case of small analyte molecules (Figure 8, right). The fractional occupancy of the sensor antibody may thus be estimated by measuring the ratio of sensor and developing antibodies that form the dualantibody "couplets." This can be readily achieved by labeling the sensor and the developing antibodies with different labels, e.g., a pair of radioactive, enzyme, or chemiluminescent markers (or even labels of entirely different nature). Fluorescent labels are potentially particularly useful in this context because, by the use of optical scanning techniques (Figure 9), they permit the scanning of arrays of antibody "microspots" distributed over a surface (each microspot directed against a different analyte), so that multiple analyte assays may be performed simultaneously on the same sample. Several



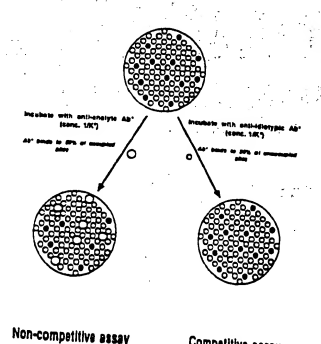


Fig. 8. Microspot immunoassay: (left) first incubation, with the fractional occupancy of antibody binding sites reflecting the analyte concentration to which the microspot has been exposed; (right) second incubation, in which the microspot is exposed to a second "developing"

Competitive assay

antibody reactive with either occupied sites (noncompetitive assay), or unoccupied sites (competitive assay) In the second incubation, a concentration of developing antibody has been selected such that only 50% of the occupied or unoccupied sites is identified

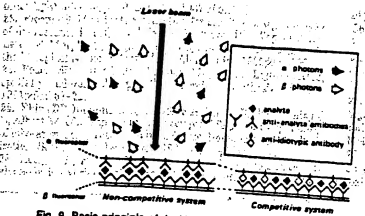


Fig. 9. Basic principle of dual-label, ambient analyte immunoassay relying on fluorescent-labeled antibodies

The ratio of a and β fluorescent photons emitted reflects the value of F (see Fig. 7) and depends solely on the analyte concentration to which the probe has been exposed. The ratio is unaffected by the amount or distribution of antibody coated (as a monomolecular layer) onto the probe surface.

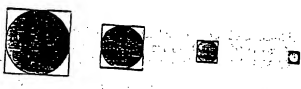
advantages stem from adopting a dual fluorescence measurement. For example, neither the amount nor the distribution of the sensor antibody within the detector's field of view is important, because the ratio of the emitted fluorescent signals is unaffected. Likewise, fluctuations in the intensity of the incident (exciting) light beam are apt to be of little significance. These advantages are additional to the basic benefit stemming from this approach, i.e., that the necessity of ensuring constancy of the amount of sensor antibody used in the assay system is removed.

Microspot Immunoassay Sensitivity

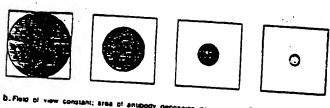
Because the microspot immunoassay methodology challenges concepts that have dominated immunoassay design theory in the past two to three decades, consideration of the potential sensitivity attainable by this approach is obviously of primary importance. The proposition that microspot assays may be at least as sensitive as conventional systems that rely on far larger amounts of antibody may readily be demonstrated by consideration of a model system. Let us postulate that sensor antibody molecules are attached to the surface of a solid support such that their binding sites remain exposed to the analyte, and that their affinity for the analyte is thereby unchanged. (The antibody concentration in the system—the number of binding sites on the support divided by the incubation volume—is unaffected by such attachment, and antibody occupancy by analyte at equilibrium will be identical to that occurring if the antibody is distributed uniformly throughout the incubation mixture.) Let us also suppose that the antibody molecules exist as a uniform monolayer of maximal surface density on the support and (to simplify discussion) are unlabeled. Then a change in the concentration of sensor antibody implies a corresponding change in the surface area over which the antibody is distributed. If, for example, the antibody affinity constant is 1011 L'mol, the total incubation volume is 1 mL, and the antibody surface density is 6000 binding sites/ μm^2 , then

a surface area of $10^5 \, \mu \mathrm{m}^2$ (i.e., 0.1 mm^2) accommodates antibody binding sites corresponding to a concentration of 0.1/K; an area of 0.01 mm² corresponds to a concentration of 0.01/K, etc. Let us further postulate that, after exposure of the sensor antibodies to a medium containing analyte at a concentration of 0.01/K (i.e., 6 × 107 molecules mL), we measure "noncompetitively" the resulting antibody occupancy (e.g., by exposure to a second, labeled, "developing" antibody directed against the analyte, forming a typical antibody sandwich). Finally, let us suppose that all occupied sites react with the developing antibody, with the latter also binding "nonspecifically" to the solid support itself at a surface density of 1 molecule/µm2.

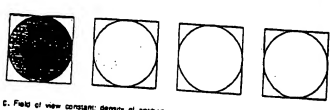
We may now consider the effects of a progressive reduction of the antibody-coated surface area from (e.g.) 1 mm² (effective antibody concentration 1/K) through 0.1 mm² (0.1/K) to 0.01 mm² (0.01/K) and below. From equation 4, the value of F for the 1 mm² area is 4.98 \times 10⁻³. Thus at equilibrium the number of analyte and labeled antibody molecules specifically bound to the area is 2.99 × 107 (i.e., about 50% of the total analyte molecules present), whereas the number of labeled antibody molecules nonspecifically bound is 10°. Thus, assuming the field of view of the detecting instrument is restricted to the area on which the sensor antibody is deposited (see Figure 10a), and (provisionally) assuming the background (or "noise") of the instrument itself to be zero (i.e., the only source of background is the non-



isto of view decreases; area of antibody deposition decreases; S/B fises



constant; area of antibody deposition decreases:



view constant; density of antibody deposition decreases; S/B falls

Fig. 10. "Capture" antibody (CAb) is assumed coated on circular (shaded) areas; the field of view of the signal-measuring instrument is represented by square (unsheded) areas (a) Reduction of both the area of deposition of CAb and the field of view results in an increase in the signal/noise (S/B) ratio. If the CAb is reduced either by reducing the antibody coated area (b) or the density of antibody coating (c) while the field of view remains unchanged, S/B falls

specifically-bound labeled antibody within the instrument's field of view), the signal/noise ratio observed for the 1 mm2 area is -30. Similarly, the value of F for a 0.1 mm² area is 9.02 × 10⁻³; the number of labeled antibody molecules specifically bound to the area is $5.41 \times$ 106, the number nonspecifically bound is 105, and the signal/noise ratio is ~54. Likewise, the signal/noise ratio for a 0.01 mm² area can be shown to be ~59. In short, the signal/noise ratio increases as the antibodycoated surface area is decreased, approaching a maximal (plateau) value of 60 as the area coated with sensor antibody falls below 0.01 mm² and tends toward zero.

If, however, a reduction in the antibody-coated area were not accompanied by a corresponding reduction in the detecting instrument's field of view, the resulting reduction in "signal" would not lead to a corresponding decrease in the background generated by nonspecifically-bound developing antibody (Figure 10b). Therefore, although reduction in the coated area would increase the fractional occupancy of the sensor antibody, the signal/noise ratio might either remain constant or fall. In these circumstances it might be advantageous to increase the coated area. Similarly, if the surface density of sensor antibody were decreased (the coated area being held constant), similar conclusions would be reached (Figure 10c).

Likewise, if the background signal generated within the detecting instrument itself (e.g., from the photocathode of a photomultiplier tube used to detect photons emitted from the antibody-coated area) were not zero, and remained constant regardless of the instrument's field of view, then a maximum signal/noise ratio would also be attained at some optimal value of the antibodycoated area, below which the ratio would fall. Because, however, one can generally reduce the size of the detector (and hence the detector-generated background) at the same rate as the size of the signal-emitting area, there is no reason—in principle—for the signal/noise ratio to diminish as the antibody-coated area is progressively reduced toward zero. Thus if we accept the signal/noise ratio as indicative of the precision of the measurement of antibody occupancy (and hence of assay sensitivity), these considerations suggest that it is advantageous to reduce the antibody-coated surface area (and, concomitantly, the sensor-antibody concentration) toward zero, although little advantage is likely to accrue from reducing the area below 0.01 mm² (and thus the antibody concentration below 0.01/K).

Were the microspot area indeed reduced to zero, both signal and noise would likewise also fall to zero (the ratio between them nevertheless remaining essentially constant), implying that no signal of any kind would, in the limit, be recorded. In practice, other statistical factors come into play when the number of individual events (e.g., photons) observed by a detecting instrument is very low, thus prohibiting a reduction of the sensor antibody concentration to zero. The point at which the reduction in the antibody-coated area causes the detectable signal to be lost sufficiently to affect the

precision of the measurement of antibody occupancy depends clearly on the specific activity of the labeled antibody used to measure the occupied binding sites; the higher the specific activity, the smaller the permissible area. Thus, given labels of very high specific activity. one can envision circumstances in which, even in a "noncompetitive" system, the optimal concentration of sensor antibody may be exceedingly low. A more general conclusion is that a variety of factors, including the characteristics of the instruments used for measuring the labeled antibody (or labeled analyte), influence immunoassay design, implying, among other things, the virtual impossibility of formulating general rules regarding this. For example, reagent concentrations that are optimal for isotopically labeled reagents used with a conventional radioisotope counter (possessing a fixed background dependent on its basic construction) are likely to be entirely different when very high-specificactivity labels are used and one has the freedom to tailor the measuring instrument to samples of any size. In short, certain conclusions based on experience of RIA and IRMA techniques may prove misleading when applied to nonisotopic methodologies, and should be viewed with caution.

A more detailed theoretical consideration of (noncompetitive) microspot immunoassay sensitivity (21) suggests that

$$C_{\min} = D^{*}_{\min} \times [(6 \times 10^{20})(1 + [Ab^{*}])]/DK[Ab^{*}]$$
 (5)

where $D = \text{surface density (binding sites/}\mu\text{m}^2\text{)}$ of sensor antibody, K = sensor antibody affinity (L/mol), $[Ab^*] =$ concentration of labeled antibody in developing solution (expressed in units of $1/K^*$, where $K^* =$ labeled antibody affinity), D* min = minimum detectable surface density of labeled antibody (molecules/ μ m²), and $C_{min} = assay$ detection limit (molecules/mL). For example, if [Ab*] = $1, D = 10^5$ molecules/ μ m², $K = 10^{11}$ L/mol, and $D^{\bullet}_{min} =$ 20 molecules/ μ m², then $C_{min} = 2.4 \times 10^6$ molecules/mL = 4×10^{-16} mol/L and the fractional occupancy of the binding sites of the sensor antibody by the minimum detectable concentration of analyte is 0.04%. Figure 11 shows the theoretical assay sensitivities attainable with use of sensor antibodies of various affinities, plotted as a function of $D*_{\min}$.

A similar theoretical analysis of competitive microspot immunoassay indicates that potential sensitivities are essentially identical to those attainable with conventional competitive methodologies. In summary, the above considerations indicate that the attainment of high microspot assay sensitivity requires close packing of molecules of sensor antibodies within the microspot area, combined with the use of an instrument capable of accurately measuring very low surface densities of developing antibodies. They also suggest that (a) microspot assay sensitivities considerably higher than those obtainable by conventional isotopically based immunoassays are achievable, and (b) if labels of very high specific activity are available, the sensitivities yielded

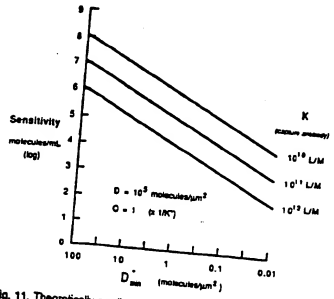


Fig. 11. Theoretically predicted sensitivity of noncompetitive microspot immunoassay plotted as a function of the minimum developing antibody density detectable within the microspot area

Postulated values of capture antibody surface density are 10s molecules/µm², and of developing antibody concentration are 1/K Currently available instruments permit detection of between 10 and 1 molecules of fluorescein-labeled

by microspot assays are unlikely to be inferior and (depending on the characteristics of the measuring instruments used) could be superior to the sensitivities achievable in macroscopic assays of conventional de-

Finally, we briefly address a further question occasionally raised in this context, i.e., the kinetic characteristics of microspot assays. Two points should be made regarding this issue. First, the smaller the microspot of sensing antibody, the lower the diffusion constraints on the velocity of the antibody/analyte binding reaction, so that at the limit (i.e., when the amount of antibody situated within the microspot area approaches zero) the kinetics of the reaction approximate those observed in a homogeneous liquid-phase system. Second, although the effective concentration of sensor antibody in the incubation medium is exceedingly low, the fractional rate at which sensor antibody binding sites within the microspot become occupied is invariably greater in this circumstance than when a relatively high concentration of antibody is used, as in conventional assays, particularly those of noncompetitive design. In other words, bearing in mind the relationship between fractional occupancy of sensor antibody and the signal/noise ratio discussed above, it is readily demonstrable that the rate at which the ratio rises is greatest when the microspot area (and the antibody contained within it) is least. Thus, given instrumentation whose field of view is restricted to the microspot area, the highest signal/noise ratio will be observed (after any selected incubation period) when the concentration of sensor antibody in the system is <0.01/K. In short, contrary perhaps to superficial impression, and to the generally accepted belief that short immunoassay incubation times require the use of very large amounts of antibody, the antibody microspot approach provides the basis of assays potentially more rapid than any currently available.

Microspot Immunoassay: Some Practical Considerations

Although various high-specific-activity antibody labels are potentially usable in this context, our preliminary studies have relied on the use of conventional fluorophors. The simultaneous measurement of dual fluorescences from small areas is, of course, well established, and the availability of improved instrumentation (e.g., the laser scanning confocal microscope), albeit not specifically designed for the present purpose, has been useful in demonstrating the feasibility of the microspot approach.

In laser scanning confocal fluorescence microscopes, a small area of the specimen is illuminated by a focused laser beam, the fluorescence photons emitted from this area being focused in turn onto a detector, typically a low-dark-current photomultiplier (22, 23). At the "confocal" point, the projection of the illumination pinhole and the back-projection of the detector pinhole coincide (Figure 12). Fluorescence photons emitted at other points thus possess a low probability of reaching the detector. Such systems contrast with conventional epifluorescence microscopes, in which the specimen is exposed to an essentially uniform flux of illumination, and yield much sharper images of fluorescent emitters situated in a defined plane of a tissue sample. Electrons spontaneously emitted by the photomultiplier photocathode contribute to the background signal of the instrument, and must-for highest microspot assay sensitivity—be minimized. Fortunately, the design of such instruments permits the photocathode to be very small in area, and this source of background can be expected

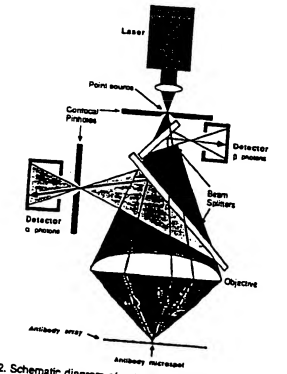


Fig. 12. Schematic diagram of a dual-ghap

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to diminish with future improvement in photomultiplier design. Other sources of background include fluorescence emitted by components in the optical system, which may not, in current instruments, have been constructed with background reduction as a prime consideration. Nevertheless, they detect with high sensitivity fluorescent signals. For example, one commercially available microscope is claimed to detect fluorescein at a density of 10 molecules/µm². Most commercially available fluorescein isothiocyanate (FITC)-labeled IgG exhibits a fluorophor/protein ratio of ~4; this implies detection limit (D*min) for antibody surface density of two or three FITC-labeled IgG molecules per micrometer2. This, in turn, implies a theoretical sensitivity for a two-site immunoassay of $\sim 2-3 \times 10^5$ analyte molecules per milliliter, assuming identical parameter values as above, or 2-3 × 104 molecules/mL if the sensing antibody has an affinity of 1012 L/mol. Clearly, sensitivity may be increased by loading more fluorophor either directly or indirectly onto the antibody.

Our preliminary studies have relied on a less sensitive microscope, albeit one possessing facilities for dualfluorescence measurement. Its argon laser emits two excitation lines at 488 and 514 nm. It is thus particularly efficient in exciting blue/green-emitting fluorophores such as FITC (excitation maximum 492 nm), but is less efficient in exciting fluorophores such as Texas Red (excitation maximum 596 nm). However, the ratiometric assay principle permits considerable variation in detection efficiencies of the two labels because the specific activities of the labeled antibody species forming the antibody couplets can be chosen to yield signal ratios approximating unity. Inefficiency of the argon laser in exciting Texas Red is thus not a major handicap in this context. Though this instrument relies on a conventional microscope and not on an optical system designed for this purpose (and thus implicitly less sensitive), it permits quantification of fluorescence signals generated from microspots of any selected area. Initial studies have revealed that, under conditions that are not optimal, the instrument is capable of detecting ~ 25 FITC-labeled and (or) 150 Texas Red-labeled IgG molecules per micrometer, while scanning an area of ~50 μm².

The development of microspot immunoassays has also necessitated closer scrutiny of the mechanisms involved in the coupling of antibodies to solid supports. In the present context, these should display a capacity to adsorb (in the form of a monolayer)—or to covalently link—a high surface density of antibody combined with low intrinsic-signal-generating properties (e.g., low intrinsic fluorescence), thus minimizing background. We have examined a number of candidate materials, such as polypropylene, Teflon®, cellulose and nitrocellulose membranes, microtiter plates (clear polystyrene plates; black, white, and clear polystyrene plates), glass slides and quartz optical fibers coated with 3-(amino propyl) triethoxy silane, etc., and several alternative protocols for achieving high monolayer coating densities. These

studies have exposed phenomena neither evident nor of importance when antibody binding to solid supports is examined at a macroscopic level. Provisionally, we have used white Dynatech Microfluor microtiter platesformulated for the detection of low fluorescence signals, and yielding high signal/noise ratios and high coating densities of functional antibodies ($\sim 5 \times 10^4$ lgG molecules/µm²)-for assay development, although such plates are not ideal. Indeed, deficiencies in the antibodydeposition methods used constitute the principal source of imprecision in assay results and the limitation in sensitivity that this implies. Clearly, this represents an area for further study and refinement of current coating techniques.

Notwithstanding the limitations of present instrumentation (which, among other things, does not permit the use of time-resolving techniques to distinguish two individual fluorescence signals either from each other or from background fluorescence) and the crudeness of present methods for coupling antibodies onto small areas; we have verified the theoretical concepts outlined above by comparing the performance of several assays when constructed in microspot format and when conventionally designed. Although unoptimized, ratiometric microspot assays have yielded sensitivity values closely approaching those of conventional optimized IRMA. As an example, the results of a ratiometric assay system for thyrotropin, with use of Texas Red- and FITC-labeled antibodies, are shown in Figure 13. Bearing in mind the well-known limitations of these and other "conventional" fluorophors when used as immunoassay reagent labels, such results are encouraging, although further work is clearly required to achieve the considerably greater sensitivity theoretically predicted with use of improved fluorophors, better antibody-microspotting techniques, and purpose-built (time-resolving) instrumentation.

The finding that highly sensitive immunoassays can be performed with far smaller amounts of antibody than

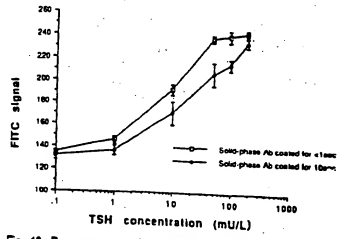


Fig. 13. Response curve in a dual-labeled microspot ratiometric assay of thyrotropin (TSH) with Texas Red-labeled solid-phase capture antibody and a developing antibody labeled with blotin/ FITC-evidin

The FITC/Texas Red ratio for each microspot was measured with a scanning contocal microscope, and plotted as a function of TSH concentration in milli-int units/L

are currently used conventionally permits in turn the construction of antibody microspot arrays enabling, in principle, the simultaneous measurement of thousands of different substances in 1-mL samples. In collaboration with investigators at the Centre for Applied Microbiological Research, Porton Down, U.K., we are presently developing various techniques for the creation of such arrays. Indeed, similar technologies have recently been used for the parallel synthesis of several different polypeptides, these enabling 10 000-microspot arrays to be constructed on silica chips approximating 1 cm2 (24). Although arrays of this capacity are unlikely to ever be required for conventional diagnostic purposes, we can anticipate that the ability to simultaneously measure many substances in the same sample will have revolutionary consequences in medicine and other similar areas. In addition, such techniques may ultimately permit the individual analysis of the multiple isoforms of certain "heterogeneous" analytes (e.g., the glycoprotein hormones), such molecular heterogeneity currently presenting a major obstacle to the standardization and interpretation of many immunological measurements (25). Moreover, although these concepts have been illustrated in an immunoassay context, they are clearly applicable to all "binding assays," including those relying on the use of DNA probes, hormone receptors, etc. For example, labeled lectins that are specific in their reactions with the sugar residues in the oligosaccharide chains of glycoprotein molecules may be used, together with specific antibodies, to impart additional "structural specificity" to sandwich assays (26, 27), possibly overcoming the limitations of antibodies per se in regard to differentiation of the glycosylation variants of the glycoprotein hormones.

Summary and Conclusion

Because of past confusion regarding the concepts of precision, sensitivity, accuracy, etc., several erroneous concepts have become incorporated within currently accepted rules of immunoassay design. In particular, much higher antibody concentrations are customarily used than are necessary to achieve very high assay sensitivity, provided that certain measurement strategies are adhered to. In this presentation, we have attempted to show that, in principle, the highest assay sensitivities are obtained by confining a small number of sensor antibody molecules onto a very small area in the form of a microspot and measuring their occupancy by an analyte, by using very high-specific-activity "developing" antibody probes, thereby maximizing the signal/noise ratio in the determination of sensor antibody occupancy. This observation, which contradicts currently accepted immunoassay design theory, in turn makes possible the measurement of an unlimited number of different analytes on a chip of very small surface area through the use of, e.g., laser scanning techniques closely analogous to those used in compact disk techniques of sound recording. Extensive experimental studies in this area, albeit conducted with relatively crude techniques and instrumentation not specifically de-

signed for these purposes, and therefore not reported in detail here, have demonstrated the feasibility of the miniaturized antibody microspot approach and the validity of the general concepts on which it is based. We are therefore confident that this represents the basis of a next-generation technology that is likely to have a revolutionary impact on all fields involving the use of binding assays. ্ত্ৰ ১ টি ৰ টিটালচে ইন্টিক্ট ডেম্ছ

References 157 to the country of the country approprie

- 1. Yalow RS, Berson SA. General principles of radioimmunosassay. In: Hayes RL, Goswitz FA, Murphy BEP, eds. Radioisotopes in medicine: in vitro studies. Oak Ridge, TN: US Atomic Energy Commission, 1968:7-39.

 2. Ekins RP, Newman B, O'Riordan JLH. Ibid.: 59-100.
- 3. Berson SA, Yalow RS. Measurement of hormones—radioimmunosssay. In: Berson SA, Yalow RS, eds. Methods in investigative and diagnostic endocrinology, Vol. 2A. Amsterdam: North Holland/Elsevier, 1973:84-135.
- 4. Ekins R, Newman B. Theoretical aspects of asturation analysis. In: Diczfalusy E, Diczfalusy A, eds. Steroid assay by protein binding. Karolinska symposia on research methods in reproductive endocrinology. Stockholm: WHO/Karolinaka, Sjukhuset,
- 5. Ekins RP. Limitations of specific activity. In Margoulies M. ed. Protein and polypeptide hormones, Part 3 (Discussions). Amsterdam: Excerpta Medica, 1968:612-6, et seq.; Ekins RP. Concentrations of tracer and antiserum, time and temperature of incubation, volume of incubation. Ibid: 672-82.
- 6. Yalow RS, Berson SA. Immunoassay of endogenous plasma insulin in man. J Clin Invest 1960;39:1157.
- 7. Ekins RP. The estimation of thyroxine in human plasms by an electrophoretic technique. Clin Chim Acta 1960;5:458-9.
- 8. Barakat RM, Ekins RP. Assay of vitamin B12 in blood-a simple method. Lancet 1961;ii:25-6.
- 9. Wide L. Bennich H., Johansson SGO. Diagnosis of allergy by an in-vitro test for allergen antibodies. Lancet 1967;ii:1105-7.
- 10. Miles LEH, Hales CN. Labeled antibodies and immunological assay systems. Nature (London) 1968;219:186-9.
- 11. Keston AS, Udenfriend S, Cannan RK. Micro-analysis of mixtures (amino acids) in the form of isotopic derivatives. J Am Chem Soc 1946;68:1390.
- 12. Avivi P, Simpson SA, Tait JF, Whitehead JK. The use of 5H and 14C-labeled acetic anhydride as analytical reagents in microbiochemistry. In: Johnston JE, Faires RA, Millett RJ, eds. Redicisotope conference, London: Butterworths, 1954:313-23.
- 13. Miles LEH, Hales CN. An immunoradiometric assay of insulin. Op. cit. (ref. 5), Part 1:61-70.
- 14. Rodbard D, Weiss GH. Mathematical theory of immunometric (labeled antibody) assay. Anal Biochem 1973;52:10-44.
- 15. Jackson TM, Marshall NJ, Ekins RP. Optimisation of immunoradiometric assays. In: Hunter WM, Corrie JET, eds. Immunoassays for clinical chemistry. Edinburgh: Churchill Livingstone,
- 16. Ekins RP. Measurement of analyte concentration. British patent no. 8 224 600, 1983.
- 17. Wide L. Solid-phase antigen-antibody systems. In: Hunter WM, Kirkham KE, eds. Radioimmunoassay methods. Edinburgh: Churchill Livingstone, 1971:405-12.
- 18. Köhler G, Milstein C. Continuous culture of fused cells secreting specific antibody of predefined specificity. Nature (London) 1975;256:495-7.
- 19. Marshall NJ, Dakubu S, Jackson T, Ekins RP. Pulsed light, time resolved fluoroimmunoassay. In: Albertini A, Ekins RP, eds. Monoclonal antibodies and developments in immunoassay. Amsterdam: Elsevier/North Holland, 1981:101-8.
- 20. Soini E, Lovgren T. Time-resolved fluorescence of lanthanide probes and applications in biotechnology [Review]. Crit Rev Anal Chem 1987:18:105-54.
- 21. Ekins RP, Chu F, Biggart E. The development of microspot, multi-analyte ratiometric immunoassay using dual fluorescentlabeled antibodies. Anal Chim Acta 1990;227:73-96.
- 22. White JG, Amos WB, Fordham M. An evaluation of confocal versus conventional imaging of biological structures by fluores-

ence light microscopy. J Cell Biol 1987;105:41-8,

23. Ploem JS. New instrumentation for sensitive image analysis of fluorescence in cells and tissues. In: Tayer DL, Waggoner AS, Lanni F, Murphy R, Birge R, eds. Applications of fluorescence in the biological sciences. New York: Alan R Lies, 1986;289-300. 24. Fodor SPA, Read JL, Pirrung MC, et al. Light-directed, spatially addressable parallel chemical synthesis. Science 1991;251:767-78.

25. Ekins RP. Immunoussay standardization. In: Kallner A, Magid E, Albert W, eds. Improvement of comparability and compatibility of laboratory assay results in life sciences. Immunoassay standardization. Scand J Clin Lab Invest 1991;51(Suppl 205):33-46.

26. Kottgen E, Hell B, Muller C, Tauber R. Demonstration of

glycosylation variants of human fibrinogen, using the new tech-

nique of glycoprotein lectin immunosorbent assay (GLIA). Biol Chem Hoppe Seyler 1988;369:1157-68.

27. Kinoshita N, Suzuki S, Matsuda Y, Taniguchi N. o-Fetoprotein antibody-lectin enzyme immunoassay to characterise sugar chains for the study of liver diseases. Clin. Chim. Acta 1989;179:143-52. 28. Shalev V, Greenberg CH, McAlpine PJ, Detection of at-

tograms of antigen by a high sensitivity enzyme-linked immunosorbent assay (HS ELISA) using a fluorogenic substrate. J Immunol Methods 1980;38:125.

29. Harris CC, Yolken RH, Kroken H, Hsu IC. Ultrasensitive enzymatic radioimmunoassay: application to detection of cholera enzymatic radioimmunoassay. apputation toxin and rotavirus. Proc Natl Acad Sci USA 1979;76:5336.

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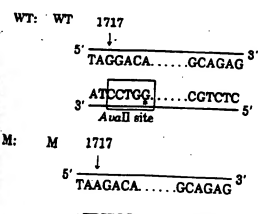
Rapid Detection of 1717-1G→A Mutation in CFTR Gene by PCR-Mediated Site-Directed Mutagenesis, Laura Cremonesi, Manuela Seia, Carmelina Magnani, and Maurizio Ferrari (1 Istituto Scientifico H.S. Raffaele, Lab. Centrale, Milano; ² Istituti Clin. di Perfezionamento, Lab. di Ricerche Clin., Milano, Italy)

Until now, among the non-AF508 mutations identified in the cystic fibrosis transmembrane conductance regulator (CFTR) gene by the Cystic Fibrosis (CF) Genetic Analysis Consortium, the ones most frequently seen in our population sample are the 1717-1G-A mutation (13/144 or 9% of the CF chromosomes) and the G542X mutation (16/190 or 8.4% of the CF chromosomes), both revealed by dot-blot hybridization of the polymerase chain reaction (PCR) product with allele-specific oligonucleotides (ASO) probes (1).

In an attempt to simplify the analysis of the most frequent mutations in the CFTR gene, we converted radiolabeled ASO detection into restriction endonuclease analysis of the amplified product.

A PCR-mediated site-directed mutagenesis (2, 3) to detect the G542X mutation by generating a novel BstNI site in the wild-type sequence had already been suggested (4).

To detect the 1717-1G→A mutation, we designed the reverse primer (5'-CTCTGCAAACTTGGAGAGGTC-3') to contain a single-base mismatch (T→G), which could create a novel AvaII restriction site [G | G(A/T)CC] in the amplified wild-type (WT) allele but not in the CF mutant (M) allele:



ATTCTGG......CGTCTC *. mutagenized base of reverse primer



Fig. 1. Detection of the 1717-1G—A mutation by PCR (b) Contact (c) Reactions were carried out with 1 μp of genomic DNA in a total volume of 100 μL containing 10 mmol/L Tris · HCI (pH 8.3), 50 mmol/L KCl, 1.6 mmol/L MgCl₂, 0.1 g/L gelatin, 200 μπο//L each of the four decay/ribonuclectide triphosphates, 2.5 units of Teq polymerase (Perion-Elmer Ceaus, Nonvalk, CT), and 100 pmoi of each of the primers. PCR conditions were as follows: denaturation at 94 °C for 1 min, annualing at 55 °C for 30 s, and extension at 72 °C for 1 min, for a total of 30 cycles. PCR products were digested for 2 h at 37 °C with 5 U of Avail and electrophoresed on 3% agarose-1% NuSieve gel for 1 h at 50 V. Bands were made visible by staining the gel with ethicium bromide. Lane 1: Heelli-digested pBR322 size marker. Lane 2: normal homozygote. Lane 3: CF patient homozygous for the 1717-1G-A mutation. Lane 4: heterozygote carrier for the 1717-1G→A mutation

For the forward primer, we used the one made available by the CF Genetic Analysis Consortium to amplify exon 11 of the CFTR gene: 5'-CAACTGTGGTTAAAGCAAT-

Digestion by AvaII enzyme of the PCR product generates two fragments of 116- and 21-bp in the wild-type alleles and leaves undigested a 137-bp fragment in the mutant alleles (Figure 1).

By combined analysis for the AF508 mutation (5) (252) 470 or 53.6% of the CF chromosomes), 1717-1G→A, and G542X, about 71% of mutations might be detected by nonisotopic analysis of the PCR product, thus allowing a faster and easier one-day procedure for carrier screening and prenatal testing.

References

- 1. Kerem B, Zielenski J, Markiewicz D, et al. Identification of mutations in regions corresponding to the two putative nucleotide (ATP)-binding folds of the cystic fibrosis gene. Proc Natl Acad Sci USA 1990;87:8447-51.
- 2. Haliassos A, Chomel JC, Baudis M, Kruh J, Kaplan JC, Kitzis A. Modification of enzymatically amplified DNA for the detection of point mutations. Nucleic Acids Res 1989;17:3606.
- 3. Friedman WE, Highamith E.Jr, Prior TW, Perry TR, Silverman LM. Cystic fibrosis deletion mutation detected by PCR-mediated site-directed mutagenesis [Tech Brief]. Clin Chem 1990;36:695-6. 4. Ng ISL, Pace R, Richard MV, et al. Methods for analysis of multiple cystic fibrosis mutations. Hum Genet (in press).

5. Ferrari M, Cremonesi L. More on detection of cystic fibrosis by polymerase chain reaction [Response to Letter]. Clin Chem 1990;36:1702_3.

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